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(54) Method and compositions for helminthic, arthropod ectoparasitic and acaridal infections with novel agents.

(57) The present invention relates to novel agents, to their production by fermentation, to methods for their recovery and concentration from crude solutions, to processes for their purification and to pharmaceutically and pharmacologically-acceptable salts thereof. Also, this invention relates to methods and compositions for the control and prevention of helminthic, arthropod ectoparasitic and acaridal infections. In warm-blooded animals, such as meat-producing animals, and poultry, by administering to said animals a therapeutically or prophylactically-effective amount of new agents designated LL-F28249 α , β , γ , δ , ϵ , ζ , η , θ , ι , κ , λ , μ , ν , and ω or mixtures thereof. The invention also relates to methods for the control of plant nematode infestations and other insecticidal activities. These novel agents are produced via a controlled condition microbiological fermentation using *Streptomyces cyaneoigriseus* ssp. *noncyanogenus*, designated LL- F28249 and having deposited accession number NRRL 15773.

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NOVEL AGENTS AND METHOD OF PRODUCTION THEREOF,
METHOD AND COMPOSITIONS FOR
HELMINTIC, ARTHROPOD ECTOPARASITIC
AND ACARIDAL INFECTIONS WITH NOVEL AGENTS

BACKGROUND OF THE INVENTION

The present invention relates to new antibiotic compounds, collectively identified as LL-F28249, which are produced by the fermentation of a nutrient medium with the strain of the microorganism Streptomyces cyaneoegriseus subsp. noncyanogenus LL-F28249, NRRL No. 15773 and to the pharmaceutically and pharmacologically-acceptable salts thereof.

5 The present invention relates to methods and compositions for preventing, treating or controlling helminthic, arthropod ectoparasitic and acaridal infections in warm-blooded animals by administering thereto an effective amount of the agents

10 (compounds) designated LL-F28249 α , β , γ , δ , ϵ , ζ , η , θ , ν κ , λ , μ , ν and ω , or mixtures thereof, such as the fermentation broth, or whole mash or the pharmaceutically and pharmacologically-acceptable salts thereof. Plant nematodes also are effectively controlled by use of these agents, mixtures and/or

15 salts. Further, these agents are effective as insecticidal agents, as well.

The diseases described above cause not only devastating effects but also serious economic problems and losses for farmers raising meat-producing animals such as swine, sheep, cattle, goats, rabbits, and poultry. Further, such diseases are a source of great concern for companion animals such as horses, dogs and cats. Although these diseases have been recognized for many years and drugs exist for the treatment

and/or prevention of such diseases, the present invention utilizes an entirely new set of active agents, isolated from a previously unknown microorganism, for the prevention, treatment or control of those diseases.

For instance, U.S. Patent 3,950,360, Aoki et al, April 5, 1976, discloses certain antibiotic substances obtained by culturing a Streptomyces microorganism, said compounds being useful as insecticides and acaracides. But as seen from the characteristics identifying such microorganism, the present microorganism is distinct, and its active components are derived from totally different microorganisms. Further, an entire series of U.S. patents relates to certain compounds produced by the fermentation of Streptomyces avermitilis, a distinct organism from the present one (U.S. Patent 4,171,314, Chabala et al, October 16, 1979; U.S. Patent 4,199,569, Chabala et al, April 22, 1980; U.S. Patent 4,206,205, Mrozik et al, June 3, 1980; U.S. Patent 4,310,519, Albers-Schonberg, January 12, 1982; U.S. Patent 4,333,925, Buhs et al, June 8, 1982). U.S. Patent 4,423,209, Mrozik, December 27, 1983 relates to the process of converting some of these less desirable components to more preferred ones. However, the present active agents identified as LL-F28249a, β , γ , δ , ϵ , ζ , η , θ , ι , κ , λ , μ , ν and ω , are derived from the fermentation of a newly discovered and previously uncultivated microorganism. Also, the present compounds and/or the fermentation broth or whole mash of microorganism Streptomyces cyaneogriseus ssp. noncyanogenus NRRL 15773, plus the pharmaceutically and pharmacologically-acceptable salts thereof (collectively referred to as active ingredient), exhibit excellent and effective treatments and/or prevention of these serious diseases of warm-blooded animals.

The full name of the microorganism LL-F28249, NRRL No. 15773, in terms of genus, species, and subspecies is Streptomyces cyaneogriseus noncyanogenus; however, for brevity it is referred to as above written throughout the specification and claims.

The strain is assigned to the genus Streptomyces based upon morphology and cell chemistry (content of the L isomer of

diaminopimelic acid). The strain's morphology and physiological data place it close to S. cyaneogriseus, as represented by ISP 5534 (ATCC 27426). Then, comparisons of the formation of gray aerial mycelium soluble pigments on media (Table A) and coiled chains of smooth conidia (3-25 spores per chain) were made. The present strain is negative for blue soluble pigment wherein the comparison strain, ISR 5534, is positive. The strains have similar reactions in the ISP carbohydrate utilization tests indicating positive for arabinose, fructose, glucose, rhamnose and xylose, while indicating negative for inositol, mannitol, raffinose and sucrose (ISP 5534 slightly positive). However, the strains differ in several characters (Table B) out of 53 in the Gordon tests. These differences support the creation of a subspecies of S. cyaneogenseus for the present microorganism.

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SUMMARY OF THE INVENTION

It is, therefore, an object of this invention to provide a novel method for the control of helminthic, arthropod ectoparasitic and acaridal infections in warm-blooded animals, particularly meat-producing animals, such as poultry, cattle, sheep, swine, rabbits, and companion animals such as horses, dogs and cats.

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It is also an object of the present invention to provide novel compositions effective for the control of said diseases in warm-blooded animals.

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It is a further object of the present invention to provide a novel method and compositions for the control of insect pests. These and further objects will become more apparent by the description of the invention.

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The culture of Streptomyces cyaneogriseus noncyanogenus (LL-F28249 and deposited under NRRL No. 15773) which produces the agents α , β , γ , δ , ϵ , ζ , η , θ , ι , κ , λ , μ , ν and ω components was isolated from mallee sand found in southern Australia.

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It has been discovered that the agents useful in the methods and compositions of the present invention are produced by the fermentation of a nutrient medium containing the strain of microorganism, Streptomyces cyaneogriseus noncyanogenus,

TABLE A

Comparison of F 28249 and ISP 5534 on ISP Morphology Test Media
 (Numbers are from NBS-ISCC)

<u>Medium</u>	<u>F 28249</u>	<u>ISP 5534</u>
Yeast-Malt (ISP 2)	A.m. ¹ Medium gray (265) V.m. Light tanish (75) Deep yellow-brown S.p. Light brown	Light to medium gray (264-265) Light tanish-white to blackish-blue (188) Light brown
Inorganic salts starch (ISP 4)	A.m. Light olive-gray (112 to medium gray (265) V.m. Deep gray to black (266-267) S.p. Grayish-yellowish- brown	Medium gray (265) Gray-purplish-blue (204) None
Glycerol- Asparagine (ISP 5)	A.m. 263 (white) to yellowish-gray (93) V.m. Black (267) to light olive brown (96) S.p. Slight brownish	263 (white) to light gray (264) Gray-purplish-blue (203-204) Light yellowish-gray
Oatmeal (ISP 3)	A.m. Yellow-gray (93) V.m. Colorless S.p. Slight yellowish	None Colorless None

1 = A.m., serial mycelium;

V.m. = vegetative mycelium;

S.p. = Soluble pigment

TABLE B

Comparison of Lederle F 28249 with ISP 5534 (Gordon Tests)

	<u>F28249</u>	<u>ISP 5534</u>
<u>Growth on/at</u>		
Salicin	+	-
10°	-	+
45°	+	-
<u>Production of</u>		
Urease	+	-
<u>Decarboxylation of</u>		
Mucate	-	+
<u>Acid Production</u>		
Raffinose	-	+
Sucrose	-	+

Both strains have the following reactions:

Positive Hydrolysis of casein, hypoxanthine, xanthine, tyrosine, adrenine, potato starch, gelatin, and esculin;
 Production of phosphatase
 Sensitivity to lysozyme
 Decarboxylation of acetate, citrate, lactate, malate, oxalate and propionate
 Acid production from arabinose, cellobiose, dextrin, fructose, galactose, glucose, glycerol, lactose, maltose, mannose, α -methyl D-glucoside, rhamnose, salicin, trehalose.

Negative Production of nitrate reductase
 Decarboxylation of benzoate and tartrate
 Acid from adonitol, dulcitol, erythritol, inositol, mannitol, sorbitol, β -methyl-D-xyloside.
 Growth on 5% NaCl

NRRL 15773. These agents include not only the fermentation broth and whole mash of said microorganism but also include the agents, LL-F29249_a, LL-F29249_B, LL-F29249_y, LL-F29249_d, LL-F29249_e, LL-F29249_z, LL-F29249_n, LL-F29249_θ, LL-F29249_l, LL-F29249_K, LL-F29249_λ, LL-F29249_μ, LL-F29249_v, and LL-F28249_w

5 The structure and stereochemistry of LL-F29249 have not been fully defined, but the proposed structures are shown below. Component LL-F29249 , is related to Hondamycin (Al-bimycin) which is disclosed in The Journal of Antibiotics, 22, No. 11, 521-526 (1969).

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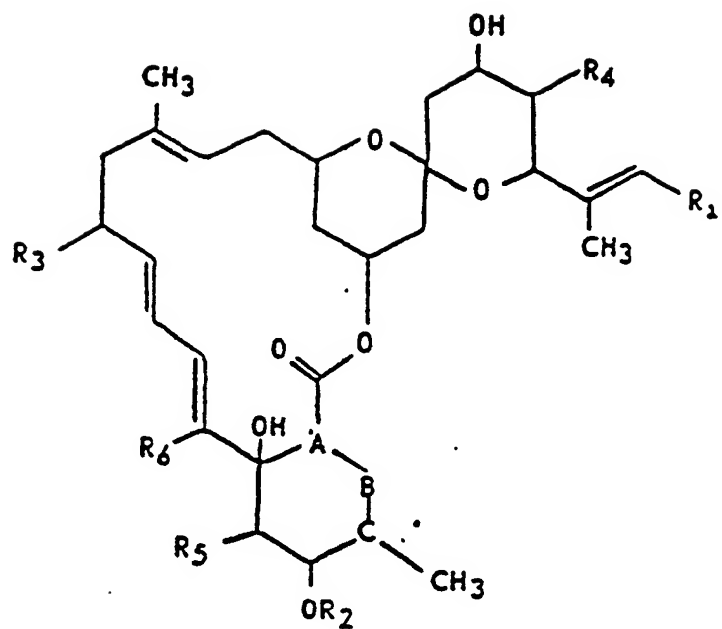
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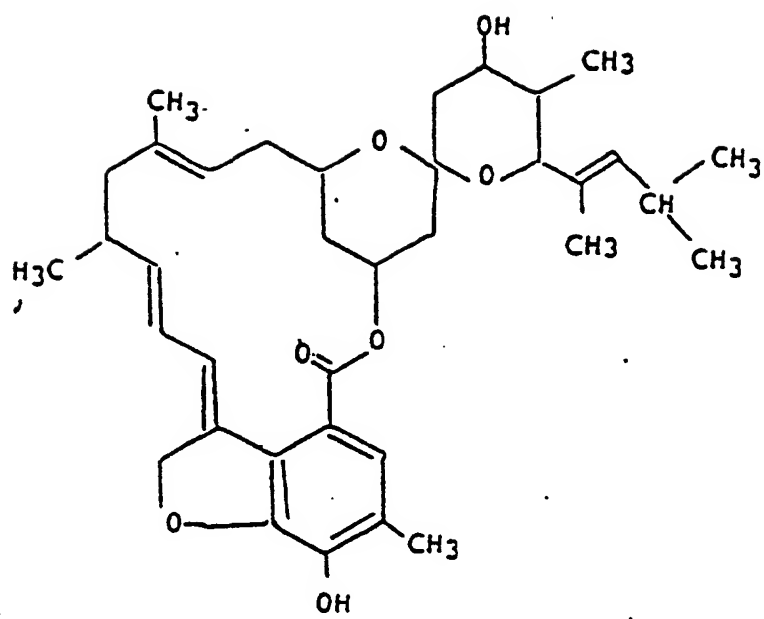


LL-F28249a-μ

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<u>Component</u>	R_1	R_2	R_3	R_4	R_5	R_6	R_5+R_6	$A-B$	$B-C$
LL-F28249 α	$CH(CH_3)_2$	H	CH_3	CH_3			-0- CH_2-	$CH-CH$	$CH=C$
LL-F28249 β	CH_3	H	CH_3	CH_3			-0- CH_2-	$CH-CH$	$CH=C$
LL-F28249 γ	CH_3	CH_3	CH_3	CH_3			-0- CH_2-	$CH-CH$	$CH=C$
LL-F28249 δ	CH_3	CH_3	CH_3	CH_3			-0- CH_2-	$CH-CH$	$CH=C$
LL-F28249 ϵ	$CH(CH_3)_2$	H	H	CH_3	CH_2OH		-0- CH_2-	$CH-CH$	$CH=C$
LL-F28249 ζ	CH_2CH_3	H	CH_3	CH_3			-0- CH_2-	$CH-CH$	$CH=C$
LL-F28249 η	$CH(CH_3)_2$	H	CH_3	CH_3			-0- CH_2-	$C=CH$	$CH=CH$
LL-F28249 θ	$CH(CH_3)_2$	H	CH_3	CH_2CH_3			-0- CH_2-	$CH-CH$	$CH=C$
LL-F28249 ι	$CH(CH_3)_2$	H	CH_2CH_3	CH_3			-0- CH_2-	$CH-CH$	$CH=C$
LL-F28249 κ	CH_3	CH_3	CH_3	H	CH_3		$CH-CH$	$CH=C$	
LL-F28249 λ	$CH(CH_3)_2$	CH_3	CH_3	CH_3	H		-0- CH_2-	$CH-CH$	$CH=C$
LL-F28249 μ	$CH(CH_3)_2$	CH_3	CH_3	H	CH_3		$CH-CH$	$CH=C$	

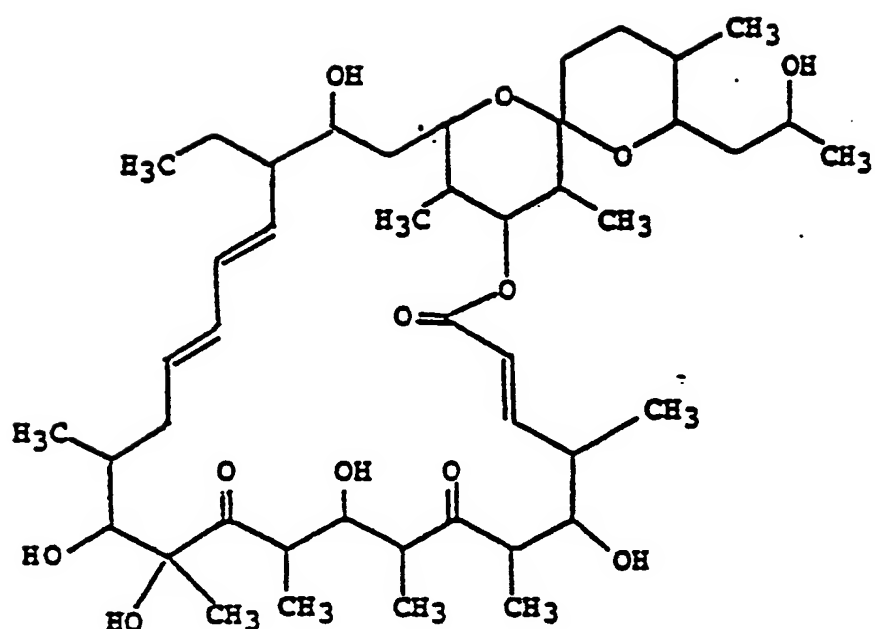
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LL-F28249v

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LL-F28249w

DESCRIPTION OF THE DRAWINGS

- FIGURE 1: Characteristic ultraviolet absorption spectrum of compound designated LL-F28249 α , NRRL 15773.
- 5 FIGURE 2: Characteristic infrared absorption spectrum of compound designated LL-F28249 α , NRRL 15773.
- FIGURE 3: Characteristic proton nuclear magnetic resonance spectrum of compound designated LL-F28249 α , NRRL 15773, in CDCl₃ solution.
- 10 FIGURE 4: Characteristic carbon-13 nuclear magnetic resonance spectrum of compound designated LL-F28249 α , NRRL 15773, in CDCl₃ solution.
- FIGURE 5: Characteristic electron impact mass spectrum of compound designated LL-F28249 α , NRRL 15773.
- 15 FIGURE 6: Characteristic ultraviolet absorption spectrum of compound designated LL-F28249 β , NRRL 15773.
- FIGURE 7: Characteristic infrared absorption spectrum of compound designated LL-F28249 β , NRRL 15773.
- 20 FIGURE 8: Characteristic proton nuclear magnetic resonance spectrum of compound designated LL-F28249 β , NRRL 15773, in CDCl₃.
- FIGURE 9: Characteristic electron impact mass spectrum of compound designated LL-F28249 β , NRRL 15773.
- 25 FIGURE 10: Characteristic ultraviolet absorption spectrum of compound designated LL-F28249 γ , NRRL 15773.
- FIGURE 11: Characteristic infrared absorption spectrum of compound LL-F28249 γ , NRRL 15773.
- 30 FIGURE 12: Characteristic proton nuclear magnetic resonance spectrum of compound LL-F28249 γ , NRRL 15773, in CDCl₃.
- FIGURE 13: Characteristic carbon-13 nuclear magnetic resonance spectrum of compound designated LL-F28249 γ , NRRL 15773, in CDCl₃.
- 35 FIGURE 14: Characteristic electron impact mass spectrum of compound designated LL-F28249 γ , NRRL 15773.
- FIGURE 15: Characteristic ultraviolet absorption spectrum of compound designated LL-F28249 ω , NRRL 15773.

- 5 FIGURE 16: Characteristic infrared absorption spectrum of compound designated LL-F28249 ω , NRRL 15773.
- 10 FIGURE 17: Characteristic proton nuclear magnetic resonance spectrum of compound designated LL-F28249 ω , NRRL 15773, in CDCl₃.
- 15 FIGURE 18: Characteristic nuclear magnetic resonance spectrum of compound designated LL-F28249 ω , NRRL 15773, in CDCl₃.
- 20 FIGURE 19: Characteristic electron impact mass spectrum of compound designated LL-F28249 ω , NRRL 15773.
- 25 FIGURE 20: Characteristic ultraviolet absorption spectrum of compound designated LL-F28249 δ , NRRL 15773.
- 30 FIGURE 21: Characteristic proton nuclear magnetic resonance spectrum of compound designated LL-F28249 δ , NRRL 15773, in CDCl₃.
- 35 FIGURE 22: Characteristic electron impact mass spectrum of compound designated LL-F28249 δ , NRRL 15773.
- FIGURE 23: Characteristic ultraviolet absorption spectrum of compound designated LL-F28249 ϵ , NRRL 15773.
- FIGURE 24: Characteristic proton nuclear magnetic resonance spectrum of compound designated LL-F28249 ϵ , NRRL 15773, in CDCl₃.
- FIGURE 25: Characteristic electron impact mass spectrum of compound designated LL-F28249 ϵ , NRRL 15773.
- FIGURE 26: Characteristic ultraviolet absorption spectrum of compound designated LL-F28249 ζ , NRRL 15773.
- FIGURE 27: Characteristic proton nuclear magnetic resonance spectrum of compound designated LL-F28249 ζ , NRRL 15773, in CDCl₃.
- FIGURE 28: Characteristic electron impact mass spectrum of compound designated LL-F28249 ζ , NRRL 15773.
- FIGURE 29: Characteristic ultraviolet absorption spectrum of compound designated LL-F28249 η , NRRL 15773.
- FIGURE 30: Characteristic proton nuclear magnetic resonance spectrum of compound designated LL-F28249 η , NRRL 15773, in CDCl₃.

- 5 FIGURE 31: Characteristic electron impact mass spectrum
 of compound designated LL-F28249 η , NRRL 15773.
- FIGURE 32: Characteristic ultraviolet absorption spectrum
 of compound designated LL-F28249 θ , NRRL 15773.
- 10 FIGURE 33: Characteristic proton nuclear magnetic reso-
 nance spectrum of compound designated LL-
 F28249 θ , NRRL 15773, in CDCl₃.
- FIGURE 34: Characteristic electron impact mass spectrum
 of compound designated LL-F28249 θ , NRRL 15773.
- 15 FIGURE 35: Characteristic ultraviolet absorption spectrum
 of compound designated LL-F28249 ι , NRRL 15773.
- FIGURE 36: Characteristic proton nuclear magnetic reso-
 nance spectrum of compound designated LL-
 F28249 ι , NRRL 15773, in CDCl₃.
- 20 FIGURE 37: Characteristic electron impact mass spectrum
 of compound designated LL-F28249 ι , NRRL 15773.
- FIGURE 38: Characteristic carbon - 13 nuclear magnetic
 resonance spectrum of compound designated LL-
 F28249 β , NRRL 15773, in CDCl₃ solution.
- 25 FIGURE 39: Characteristic ultraviolet absorption spectrum
 of compound designated LL-F28249 κ , NRRL 15773.
- FIGURE 40: Characteristic infrared absorption spectrum of
 compound designated LL-F28249 κ , NRRL 15773.
- FIGURE 41: Characteristic electron impact mass spectrum
 of compound designated LL-F28249 κ , NRRL 15773.
- 30 FIGURE 42: Characteristic proton nuclear magnetic reso-
 nance spectrum of compound designated LL-
 F28249 κ , NRRL 15773.
- FIGURE 43: Characteristic carbon - 13 nuclear magnetic
 resonance spectrum of compound designated LL-
 F28249 κ , NRRL 15773.
- 35 FIGURE 44: Characteristic ultraviolet absorption spectrum
 of compound designated LL-F28249 λ , NRRL 15773.
- FIGURE 45: Characteristic infrared absorption spectrum of
 compound designated LL-F28249 λ , NRRL 15773.
- FIGURE 46: Characteristic electron impact mass spectrum
 of compound designated LL-F28249 λ , NRRL 15773.

FIGURE 47: Characteristic proton nuclear magnetic resonance spectrum of compound designated LL-F28249 λ , NRRL 15773.

5 FIGURE 48: Characteristic carbon - 13 nuclear magnetic resonance spectrum of compound designated LL-F28249 λ , NRRL 15773.

FIGURE 49: Characteristic ultraviolet absorption spectrum of compound designated LL-F28249 μ , NRRL 15773.

10 FIGURE 50: Characteristic infrared absorption spectrum of compound designated LL-F28249 μ , NRRL 15773.

FIGURE 51: Characteristic electron impact mass spectrum of compound designated LL-F28249 μ , NRRL 15773.

15 FIGURE 52: Characteristic proton nuclear magnetic resonance spectrum of compound designated LL-F28249 μ , NRRL 15773.

FIGURE 53: Characteristic ultraviolet absorption spectrum of compound designated LL-F28249 ν , NRRL 15773.

FIGURE 54: Characteristic infrared absorption spectrum of compound designated LL-F28249 ν , NRRL 15773.

20 FIGURE 55: Characteristic electron impact mass spectrum of compound designated LL-F28249 ν , NRRL 15773.

FIGURE 56: Characteristic proton nuclear magnetic resonance spectrum of compound designated LL-F28249 ν , NRRL 15773.

25 FIGURE 57: Characteristic carbon - 13 nuclear magnetic resonance spectrum of compound designated LL-F28249 ν , NRRL 15773.

DETAILED DESCRIPTION OF THE INVENTION

It has been discovered that the above-mentioned agents, as well as the fermentation broth and whole mash of said microorganism, are especially effective for controlling helminthic, arthropod ectoparasitic and acaridal infections in meat-producing animals such as cattle, sheep, swine, rabbits, poultry, such as chickens, turkeys, ducks, geese, quail, and pheasants and companion animals.

In practice, the present invention involves the method of preventing, controlling or treating said infections, in warm-blooded animals by administering orally, parentally, or topically thereto, a prophylactically, pharmaceutically or therapeutically-effective amount of the fermentation broth or whole mash of microorganism Streptomyces cyaneoegriseus noncyanogenus, NRRL 15773, the fermentation broth or whole mash of said microorganism containing compounds designated LL-F28249 α , β , γ , δ , ϵ , ζ , η , θ , ι , κ , λ , μ , ν and ω , compounds designated as LL-F28249 α , LL-F28249 β , LL-F28249 γ , LL-F28249 δ , LL-F28249 ϵ , LL-F28249 ζ , LL-F28249 η , LL-F28249 θ , LL-F28249 ι , LL-F28249 κ , LL-F28249 λ , LL-F28249 μ , LL-F28249 ν , and LL-F28249 ω , as identified and characterized herein, or the pharmaceutically and pharmacologically-acceptable salts thereof (collectively referred to as active ingredient).

Although administration of the compound or fermentation broth/whole mash (hereinafter broth or mash) will generally be most practical in or with the feed or in the drinking water, the above-said compounds, broth or mash, or pharmaceutically and pharmacologically-acceptable salts thereof, may also be administered to individual hosts in the form of tablets, drenches, gels, capsules, or the like, or by injection in the form of a paste, gel, pellet, or solution. These latter methods of administration are, of course, less practical for the treatment of large groups of animals, but they are quite

practical for use on a small scale or on an individual basis.

When the agents (antibiotics) LL-F28249 α , β , γ , δ , ϵ , ζ , η , θ , ι , κ , λ , μ , ν or ω or the fermentation broth or whole mash of Streptomyces cyaneoaristatus noncyanogenus NRRL 15773 are used as prophylactic or therapeutic treatments of helminthic, arthropod ectoparasitic and acaridal infections, in animals and poultry, generally about 0.05 ppm to 500.0 ppm, and preferably 0.1 ppm to 300 ppm of the agent or broth or mash above-described, administered in the diet or drinking water of the animal, is effective for preventing, controlling, or treating said infections in those animals.

Medicated feeds useful in the method of the present invention are usually prepared by thoroughly admixing about 0.00001% by weight to about 0.01% by weight of the agent (antibiotic) or above-described broth or mash with a nutritionally-balanced feed, as for example, the feed described in the examples hereinafter.

When using the compounds and/or broth or mash of the present invention for the prevention or control of helminths, arthropod ectoparasites and acarides, the active agent is generally first prepared as an animal feed premix. The premix usually contains a relatively high percentage of the active ingredient and is generally blended with the animal's feed just prior to administration. If desired, the feed premix may also be applied as a top dressing for the animal's daily ration.

Feed premixes or concentrates, useful in the practice of the present invention, may be prepared by admixing about 0.1% to 5.0% by weight of the above-identified agents, broth or mash, or pharmaceutically and pharmacologically-acceptable salts thereof, with about 99.9% to 95% by weight of a suitable carrier or diluent.

Carriers suitable for use to make up the feed supplement compositions include the following: alfalfa meal, soybean meal, cottonseed oil meal, linseed oil meal, sodium chloride, calcium carbonate, calcium sulfate, cornmeal, cane molasses, urea, bone meal, corncob meal, rice hull meal, and the like. The carrier promotes an essentially uniform distribution of the active ingredient in the finished feed into which the supplement is blended. It thus performs an important function by ensuring proper distribution of the active ingredient, i.e., about 0.1 ppm to 100 ppm thereof, throughout the feed. This is equivalent to 0.00001% to 0.01%, by weight, of the active ingredient in the finished feed. In practice, usually one or more pounds of premix is added per ton of feed to obtain the desired level of agent (antibiotic) or broth or mash in the finished feed.

If the supplement or premix is used as a top dressing for feed, it likewise helps to ensure uniformity of distribution of the active ingredient across the top of the dressed feed.

Since the compounds of this invention and their pharmaceutically and pharmacologically-acceptable salts are relatively insoluble in water, it is generally desirable, when administering any such compound in the animal's drinking water, to dissolve the active ingredient in an organic solvent such as methanol, ethanol, acetone, DMSO, oleic acid, linoleic acid, propylene glycol, or the like, and admix with the solution a small amount of surfactant and/or dispersing agent to assure solution and/or dispersion of the active ingredient in the animal's drinking water.

Advantageously, where the treatment of a small number of the larger meat-producing animals is required to control parasitic infection therein, the agents LL-F28249 α , β , γ , δ , ϵ , ζ , η , θ , ι , κ , λ , μ , ν and ω , broth or mash, or pharmaceutically or pharmacologically-acceptable salts thereof may be

orally administered, on a daily basis, to the host animal in the form of a medicated gel.

The active ingredients of the invention have also exhibited nematocidal activity against plant nematodes as demonstrated by effectiveness in controlling the free living soil nematode, C. elegans. Compositions containing these active ingredients for controlling plant nematodes can be formulated into either liquids or wettable powders. Liquid compositions include about 5% to 20%, w/w, of the active ingredient (active agent, fermentation broth, whole mash or salts) with appropriate amounts of a solvent such as methanol, ethanol, acetone, acetonitrile, and others, and the remainder water. Wettable powders include about 5% to 20%, w/w, of the active ingredient, about 1% to 10% of surfactant, and inert carriers, such as clays, vermiculite, carbon black or the like. About 0.1 to 1.4 kg per hectare is applied to the foliage of plants, the soil in which they are grown or into the trunks thereof.

These agents also are active as topical insecticides, stomach poisons and systemic insecticides and are especially effective for controlling insects of the orders Lepidoptera, Coleoptera, Homoptera, Diptera and Thysanoptera. Plant mites, acarids, additionally are controlled by the agents of the present invention.

These agents generally are applied as dilute, solid or liquid compositions to the breeding ground, food supply or habitat of such insects and/or acarids. The rate of application to such loci include about 0.01 kg/ha to about 8.0 kg/ha, preferably about 0.05 kg/ha to about 0.5 kg/ha.

Surfactants useful in wettable powders of the present invention include those commonly used for formulations of such wettable powders, preferably alkylbenzene sulfonate sodium salts. Bentonite, clay or mixtures thereof are preferred carriers.

Additionally, the active ingredients of the invention also have demonstrated systemic insecticidal activity against m. ovinus in sheep.

5 In practice, generally about 0.02mg/kg/day to about 3.0 mg/kg/day is effective for controlling parasitic infections in cattle, sheep, and swine and companion animals. For prolonged use, rates as low as 0.002 mg/kg of body weight/day may be employed.

10 Also in practice, about 0.1 mg per kg to 100 mg per kg is administered to animals infected with helminths.

The physiochemical characteristics for the α , β , γ , δ , ϵ , ζ , η , θ , ι , κ , λ , μ , ν and ω components of LL-F28249 are described below:

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DETAILED DESCRIPTION OF THE INVENTION

The physiochemical characteristics for the α , β , γ , δ , ϵ , ζ , η , θ , κ , λ , μ , ν and ω components of LL-F28249 are described below:

LL-F28249_g:

- 5 1) Molecular weight: 612 (FAB-MS);
- 2) Molecular formula: C₃₆H₅₂O₈;
- 3) Specific optical rotation: $[\alpha]_D^{26} = +133 \pm 3^\circ$ (C 0.3, acetone);
- 4) Ultraviolet absorption spectrum: as shown in Figure I UV_{MAX}^{CH₃OH} = 244 nm (ϵ 28,000);
- 10 5) Infrared absorption spectrum: as shown in Figure II (KBr disc): 3439, 2960, 2925, 1714, 1454, 1374, 1338, 1171, 1120, 996, 967 cm⁻¹;
- 15 6) Proton nuclear magnetic resonance spectrum (CDCl₃): as shown in Figure III;
- 7) Carbon-13 nuclear magnetic resonance spectrum (CDCl₃): as shown in Figure IV and described in Table I; and
- 18 8) Electron impact mass spectrum: as shown in Figure V with accurate mass measurements and proposed elemental compositions indicated in Table II.
- 20

LL-F28249_B:

- 1) Molecular weight: 584 (FAB-MS);
- 2) Molecular formula: C₃₄H₄₈O₈;
- 25 3) Specific optical rotation: $[\alpha]_D^{26} = +125^\circ$ (C 0.30 acetone).
- 4) Ultraviolet absorption spectrum: as shown in Figure VI UV_{MAX}^{CH₃OH} = 244 nm (ϵ 25,600);
- 5) Infrared absorption spectrum: as shown in Figure VII (KBr disc): 3520, 2910, 1735, 1717, 1450, 1375, 1335, 1180, 1170, 1119, 993, 727 cm⁻¹;
- 30 6) Proton nuclear magnetic resonance spectrum (CDCl₃): as shown in Figure VIII;
- 7) Carbon-13 nuclear magnetic resonance spectrum (CDCl₃): as shown in Figure XXXVIII and described in Table II A; and
- 36 8) Electron impact mass spectrum: as shown in Figure IX with accurate mass measurements and proposed ele-

mental compositions indicated in Table III.

LL-F28249_y:

- 1) Molecular weight: 598 (FAB-MS);
- 2) Molecular formula: C₃₅H₅₀O₈;
- 3) Specific optical rotation: $[\alpha]_D^{26} = +150 \pm 4^\circ$ (C 0.3,
acetone);
- 4) Ultraviolet absorption spectrum: as shown in Figure
X UV_{MAX}^{CH₃OH} = 244 nm (ϵ 27,100);
- 5) Infrared absorption spectrum: as shown in Figure XI
(KBr disc): 3510, 2910, 1735, 1715, 1452, 1375,
1338, 1182, 1172, 1119, 995 cm⁻¹;
- 10 6) Proton nuclear magnetic resonance spectrum (CDCl₃):
as shown in Figure XII;
- 7) Carbon-13 nuclear magnetic resonance spectrum
(CDCl₃): as shown in Figure XIII and described in
Table IV; and
- 15 8) Electron impact mass spectrum: as shown in Figure
XIV with accurate mass measurements and proposed
elemental compositions indicated in Table V.

LL-F28249_w:

- 20 1) Molecular weight: 806 (FAB-MS);
- 2) Molecular formula: C₄₅H₇₄O₁₂;
- 3) Specific optical rotation: $[\alpha]_D^{26} = -49 \pm 3^\circ$ (C 0.35,
methanol);
- 4) Ultraviolet absorption spectrum: as shown in
Figure XV UV_{MAX}^{CH₃OH} = 225 nm (ϵ 27,400)
232 nm (ϵ 25,700);
- 25 5) Infrared absorption spectrum: as shown in Figure XVI
(KBr disc): 3480, 2965, 2935, 2880, 1703, 1647,
1458, 1380, 1292, 1223, 1135, 1098, 984 cm⁻¹;
- 30 6) Proton nuclear magnetic resonance spectrum (CDCl₃):
as shown in Figure XVII;
- 7) Carbon-13 nuclear magnetic resonance spectrum
(CDCl₃): as shown in Figure XVIII and described in
Table VI; and
- 35 8) Electron impact mass spectrum: as shown in Figure
XIX with accurate mass measurements and proposed
elemental compositions indicated in Table VII.

LL-F28249₆:

- 1) Molecular weight: 616 (EI-MS)
- 2) Molecular formula: C₃₅H₅₂O₉
- 3) HPLC retention volume of 14.0 ml in the system indicated in Table VIII;
- 5 4) Ultraviolet absorption spectrum (methanol): as shown in Figure XX;
- 5) Proton nuclear magnetic resonance spectrum (CDCl₃): as shown in Figure XXI; and
- 6) Electron impact mass spectrum: as shown in Figure
- 10 XXII.

LL-F28249_E:

- 1) Molecular weight: 598 (EI-MS)
- 2) Molecular formula: C₃₅H₅₀O₈
- 3) HPLC retention volume of 14.8 ml in the system indicated in Table VIII;
- 15 4) Ultraviolet absorption spectrum (methanol): as shown in Figure XXIII;
- 5) Proton nuclear magnetic resonance spectrum (CDCl₃): as shown in Figure XXIV; and
- 20 6) Electron impact mass spectrum: as shown in Figure XXV.

LL-F28249_C:

- 1) Molecular weight: 598 (EI-MS)
- 2) Molecular formula: C₃₅H₅₀O₈
- 25 3) HPLC retention volume of 16.0 ml in the system indicated in Table VIII;
- 4) Ultraviolet absorption spectrum (methanol): as shown in Figure XXVI;
- 5) Proton nuclear magnetic resonance spectrum (CDCl₃):
- 30 as shown in Figure XXVII; and
- 6) Electron impact mass spectrum: as shown in Figure XXVIII.

LL-F28249_H:

- 1) Molecular weight: 612 (EI-MS)
- 35 2) Molecular formula: C₃₆H₅₂O₈
- 3) HPLC retention volume of 23.5 ml in the system indicated in Table VIII;

- 4) Ultraviolet absorption spectrum (methanol): as shown in Figure XXIX;
5) Proton nuclear magnetic resonance spectrum (CDCl_3): as shown in Figure XXX; and
6) Electron impact mass spectrum: as shown in Figure
5 XXXI.

LL-F28249_a:

- 1) Molecular weight: 626 (EI-MS)
2) Molecular formula: $\text{C}_{37}\text{H}_{54}\text{O}_8$
3) HPLC retention volume of 24.5 ml in the system indicated in Table VIII;
10 4) Ultraviolet absorption spectrum (methanol): as shown in Figure XXXII;
5) Proton nuclear magnetic resonance spectrum (CDCl_3): as shown in Figure XXXIII; and
15 6) Electron impact mass spectrum: as shown in Figure XXXIV.

LL-F28249_L:

- 1) Molecular weight: 626 (EI-MS)
2) Molecular formula: $\text{C}_{37}\text{H}_{54}\text{O}_8$
20 3) HPLC retention volume of 26.0 ml in the system indicated in Table VIII;
4) Ultraviolet absorption spectrum (methanol): as shown in Figure XXXV;
25 5) Proton nuclear magnetic resonance spectrum (CDCl_3): as shown in Figure XXXVI; and
6) Electron impact mass spectrum: as shown in Figure XXXVII.

LL-F28249_k:

- 30 1) Molecular weight: 584 (EI-MS);
2) Molecular formula: $\text{C}_{35}\text{H}_{52}\text{O}_7$;
3) Specific optical rotation: $[\alpha]^{26}_{\text{D}} = +189^{\circ}$ - (C 0.165 acetone);
4) Ultraviolet absorption spectrum: as shown in Figure XXXIX UV CH_3OH $_{\text{MAX}}^{=241\text{nm}}$ (E20,400);
35 5) Infrared absorption spectrum: as shown in Figure XL (KBr disc);

- 6) Electron impact mass spectrum: as shown in Figure XLI;
7) Proton nuclear magnetic resonance spectrum (CDCl_3);
as shown in Figure XLII; and
8) Carbon-13 nuclear magnetic resonance spectrum (CDCl_3);
5 as shown in Figure XLIII and described in Table IX.

LL-F28249_λ:

- 1) Molecular weight: 626 (FAB-MS);
2) Molecular formula: $\text{C}_{37}\text{H}_{54}\text{O}_8$;
3) Specific optical rotation: $[\alpha]_D^{26} = +145^\circ$ (C, 0.23
10 acetone);
4) Ultraviolet absorption spectrum: as shown in Figure
XLIV UV CH_3OH MAX = 244 nm (E30,000);
5) Infrared absorption spectrum: as shown in Figure XLV
(KBr disc);
15 6) Electron impact mass spectrum: as shown in Figure
XLVI;
7) Proton nuclear magnetic resonance spectrum (CDCl_3);
as shown in Figure XLVII; and
8) Carbon-13 nuclear magnetic resonance spectrum (CDCl_3);
20 as shown in Figure XLVIII and described in Table X.

LL-F28249_μ:

- 1) Molecular weight: 612 (EI-MS);
2) Molecular formula: $\text{C}_{37}\text{H}_{56}\text{O}_7$;
3) Ultraviolet absorption spectrum: as shown
25 in Figure XLIX UV CH_3OH MAX = 241 nm (E16,800);
4) Infrared absorption spectrum: as shown in Figure L
(KBr disc);
5) Electron impact mass spectrum: as shown in Figure
LI;
30 6) Proton nuclear magnetic resonance spectrum (CDCl_3);
as shown in Figure LII.

LL-F28249_ν:

- 1) Molecular weight: 592 (EI-MS);
2) Molecular formula: $\text{C}_{36}\text{H}_{48}\text{O}_7$;
35 3) Specific optical rotation: $[\alpha]_D^{26} +131^\circ$ - (C .325,

- acetone);
- 4) Ultraviolet absorption spectrum: as shown in Figure LIII UV CH_3OH MAX = 256 (E20,500); 358(E 8,830);
- 5) Infrared absorption spectrum: as shown in Figure LIV (KBr disc);
- 5 6) Electron impact mass spectrum: as shown in Figure LV;
- 7) Proton nuclear magnetic resonance spectrum (CDCl_3); as shown in Figure LVI; and
- 8) Carbon-13 nuclear magnetic resonance spectrum (CDCl_3); as shown in Figure LVII, and described in Table XI.
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TABLE I
Carbon-13 NMR Data for LL-F28249a

Carbon	Chemical Shift (ppm)	Proton Substitution	Carbon	Chemical Shift (ppm)	Proton Substitution
1	173.4	q ²	18	67.8	CH
2	142.8	CH	19	67.7	CH
3	139.4	q	20	48.4	CH ₂
4	137.7	q	21	45.7	CH
5	137.3	q	22	41.1	CH ₂
6	137.2	CH	23	40.7	CH ₂
7	130.6	q	24	36.1	CH ₂
8	123.3	CH	25	36.0	CH
9	120.3 ³	CH	26	35.9	CH
10	118.0	CH	27	34.7	CH ₂
11	99.7	q	28	26.8	CH
12	80.2	q	29	22.8 ⁴	CH ₃
13	79.3	CH	30	22.2	CH ₃
14	76.7	CH	31	19.9	CH ₃
15	69.3	CH	32	15.5	CH ₃
16	68.5	CH	33	13.9	CH ₃
17	68.4	CH ₂	34	11.0	CH ₃

¹ Downfield from TMS; CDCl₃ solution.

² q = quaternary carbon.

^{3,4} Two unresolved signals.

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TABLE II
High Resolution Mass Measurements
for LL-F28249a

<u>m/z</u>	<u>Elemental Composition</u>
612.3705	C ₃₆ H ₅₂ O ₈
594.3543	C ₃₆ H ₅₀ O ₇
576.3472	C ₃₆ H ₄₈ O ₆
484.3211	C ₃₀ H ₄₄ O ₅
482.2648	C ₂₉ H ₃₈ O ₆
466.3097	C ₃₀ H ₄₂ O ₄
448.2987	C ₃₀ H ₄₀ O ₃
442.2375	C ₂₆ H ₃₄ O ₆
425.2327	C ₂₆ H ₃₃ O ₅
354.2181	C ₂₃ H ₃₀ O ₃
314.1877	C ₂₀ H ₂₆ O ₃
278.1144	C ₁₅ H ₁₈ O ₅
265.1786	C ₁₆ H ₂₅ O ₃
248.1405	C ₁₅ H ₂₀ O ₃
247.1705	C ₁₆ H ₂₃ O ₂
237.1838	C ₁₅ H ₂₅ O ₂
219.1740	C ₁₅ H ₂₃ O
151.0753	C ₉ H ₁₁ O ₂

TABLE IIaCarbon-13 NMR Data for LL-F282498

<u>Carbon</u>	<u>Chemical Shift(ppm)</u> *	<u>Carbon</u>	<u>Chemical Shift(ppm)</u>
1	173.3	18	68.3
2	142.6	19	67.8
3	139.5	20	67.7
4	137.7	21	48.4
5	137.3	22	45.7
6	133.9	23	41.0
7	123.8	24	40.8
8	123.4	25	36.1
9	120.3	26	35.9 **
10	120.2	27	34.7
11	118.0	28	22.3
12	99.7	29	19.8
13	80.2	30	15.5
14	79.4	31	13.8
15	76.7	32	13.1
16	69.2	33	10.8
17	68.6		

* Downfield from TMS; CDCl₃ solution

** Two unresolved signals

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TABLE III
High Resolution Mass Measurements
for LL-F282498

<u>m/z</u>	<u>Elemental Composition</u>
584.3388	C ₃₄ H ₄₈ O ₈
566.3306	C ₃₄ H ₄₆ O ₇
456.2864	C ₂₈ H ₄₀ O ₅
442.2391	C ₂₆ H ₃₄ O ₆
438.2780	C ₂₈ H ₃₈ O ₄
425.2331	C ₂₆ H ₃₃ O ₅
354.2187	C ₂₃ H ₃₀ O ₃
314.1858	C ₂₀ H ₂₆ O ₃
278.1168	C ₁₅ H ₁₈ O ₅
237.1491	C ₁₄ H ₂₁ O ₃
219.1380	C ₁₄ H ₁₉ O ₂
209.1534	C ₁₃ H ₂₁ O ₂
191.1418	C ₁₃ H ₁₉ O
151.0750	C ₉ H ₁₁ O ₂

TABLE IV
Carbon-13 NMR Data for LL-F28249 Y

<u>Carbon</u>	<u>Chemical Shift¹</u> (ppm)	<u>Carbon</u>	<u>Chemical Shift</u> (ppm)
1	173.6	19	68.3
2	142.4	20	67.9
3	139.9	21	57.7
4	137.3	22	48.5
5	136.0	23	45.8
6	134.0	24	41.2
7	123.8	25	40.8
8	123.6	26	36.2
9	120.4	27	36.1
10	119.6	28	36.0
11	118.5	29	34.8
12	99.8	30	22.3
13	80.5	31	19.9
14	77.8	32	15.5
15	77.0	33	13.8
16	76.8	34	13.1
17	69.3	35	10.8
18	68.6		

¹Downfield from TMS; CDCl₃ solution.

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TABLE V
High Resolution Mass Measurements
for LL-F28249

<u>m/z</u>	<u>Elemental Composition</u>
598.3543	C ₃₅ H ₅₀ O ₈
580.3422	C ₃₅ H ₄₈ O ₇
562.3292	C ₃₅ H ₄₆ O ₆
496.2824	C ₃₀ H ₄₀ O ₆
484.2440	C ₂₈ H ₃₆ O ₇
478.2687	C ₃₀ H ₃₈ O ₅
456.2576	C ₂₇ H ₃₆ O ₆
438.2772	C ₂₈ H ₃₈ O ₄
425.2341	C ₂₆ H ₃₃ O ₅
420.2651	C ₂₈ H ₃₆ O ₃
354.2199	C ₂₃ H ₃₀ O ₃
314.1875	C ₂₀ H ₂₆ O ₃
292.1307	C ₁₆ H ₂₀ O ₅
288.2075	C ₁₉ H ₂₈ O ₂
248.1397	C ₁₅ H ₂₀ O ₃
237.1490	C ₁₄ H ₂₁ O ₃
219.1382	C ₁₄ H ₁₉ O ₂
209.1544	C ₁₃ H ₂₁ O ₂
191.1435	C ₁₃ H ₁₉ O
151.0759	C ₉ H ₁₁ O ₂

TABLE VI

<u>Carbon</u>	<u>Chemical Shift¹</u> (ppm)	<u>Carbon</u>	<u>Chemical Shift</u> (ppm)
1	220.7	23	42.2 ²
2	219.6	24	40.4
3	165.2	25	38.3
4	148.7	26	37.6
5	133.1	27	36.1
6	132.3	28	34.8
7	132.1	29	33.5
8	130.2	30	30.1
9	122.3	31	26.6
10	100.0	32	25.4
11	82.9	33	24.5
12	75.9	34	23.0
13	73.0	35	21.1
14	72.7	36	17.9
15	72.6	37	14.3
16	72.1	38	14.2
17	69.0	39	12.1
18	67.3	40	11.5
19	63.6	41	10.9
20	51.4	42	8.7
21	46.2	43	8.3
22	45.7	44	5.7

¹Downfield from TMS; CDCl₃ solution.

Two unresolved signals.

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TABLE VII
High Resolution Mass Measurements
for LL-F28249 w

<u>m/z</u>	<u>Elemental Composition</u>
462.3350	C ₂₈ H ₄₆ O ₅
444.3237	C ₂₈ H ₄₄ O ₄
425.2534	C ₂₃ H ₃₇ O ₇
407.2439	C ₂₃ H ₃₅ O ₆
406.3046	C ₂₅ H ₄₂ O ₄
387.2895	C ₂₅ H ₃₉ O ₃
337.2010	C ₁₉ H ₂₉ O ₅
297.2031	C ₁₇ H ₂₉ O ₄
279.1944	C ₁₇ H ₂₇ O ₃
261.1851	C ₁₇ H ₂₅ O ₂
253.1797	C ₁₅ H ₂₅ O ₃
235.1697	C ₁₅ H ₂₃ O ₂
224.1754	C ₁₄ H ₂₄ O ₂
209.1530	C ₁₃ H ₂₁ O ₂
207.1744	C ₁₄ H ₂₃ O
184.1458	C ₁₁ H ₂₀ O ₂
179.1048	C ₁₁ H ₁₅ O ₂
173.1205	C ₉ H ₁₇ O ₃
167.1051	C ₁₀ H ₁₅ O ₂
155.1069	C ₉ H ₁₅ O ₂

<u>Compound</u>	<u>Retention Volume*(ml)</u>
LL-F28249 α	19.8
LL-F28249 δ	14.0
LL-F28249 ϵ	14.8
LL-F28249 ζ	16.0
LL-F28249 η	23.5
LL-F28249 θ	24.5
LL-F28249 ι	26.0

*System includes a column 3.9mm x 30cm packed with C₁₈ reverse phase packing developed with methanol:water (80:20) at 1.0 ml/minute, detection was by absorbance at 254 nm.

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TABLE IX
Carbon-13 NMR Data for LL-F28249_K

<u>Carbon</u>	<u>Chemical Shift(ppm)*</u>	<u>Carbon</u>	<u>Chemical Shift(ppm)</u>
1	173.9	19	56.7
2	140.7	20	48.4
3	138.3	21	47.7
4	136.6	22	41.1
5	136.5	23	40.6
6	133.8	14	37.1
7	124.7	25	36.3
8	124.4	26	36.0
9	123.8	27	35.9
10	120.1	28	34.6
11	118.5	29	22.0
12	99.7	30	19.3
13	77.2	31	16.0
14	76.6**	32	13.8
15	76.5	33	13.3
16	69.3	34	13.1
17	68.6	35	10.7
18	67.3		

* Downfield from TMS; CDCl₃ solution.

** Coincident with CDCl₃ signals.

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TABLE XCarbon-13 NMR Data for LL-F28249λ

<u>Carbon</u>	<u>Chemical Shift(ppm)*</u>	<u>Carbon</u>	<u>Chemical Shift(ppm)</u>
1	173.6	19	68.3
2	142.5	20	67.9
3	139.8	21	57.8
4	137.4	22	48.6
5	137.2	23	45.8
6	136.0	24	41.2
7	130.7	25	40.9
8	123.6	26	36.1 **
9	120.3	27	36.0
10	119.7	28	34.9
11	118.6	29	26.9
12	99.8	30	23.0 **
13	80.5	31	22.4
14	77.7	32	20.0
15	77.6	33	15.7
16	76.7	34	14.0
17	69.3	35	11.1
18	68.6		

* Downfield from TMS; CDCl₃ solution.

** Two unresolved signals.

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TABLE XICarbon-13 NMR Data for LL-F28249v

<u>Carbon</u>	<u>Chemical Shift(ppm)*</u>	<u>Carbon</u>	<u>Chemical Shift(ppm)</u>
1	167.4	18	69.4
2	150.5	19	68.7
3	142.9	20	68.3
4	142.0	21	48.4
5	137.2 **	22	41.0 **
6	132.1	23	35.9
7	130.7	24	35.6
8	125.8	25	35.5
9	125.5	26	34.4
10	124.2	27	29.7
11	123.7	28	26.8
12	123.2	29	22.9
13	121.3	30	22.8
14	118.0	31	22.1
15	100.0	32	15.3
16	76.7	33	13.9
17	74.6	34	11.0

* Downfield from TMS; CDCl₃ solution.

** Two unresolved signals.

TABLE XII
Chromatographic Data

<u>Component</u>	<u>TLC *</u> <u>Relative Rf</u>	<u>HPLC **</u> <u>Retention Time (minutes)</u>
a	1.00	13.8
b	.797	9.3
γ	1.42	12.6
δ	.758	10.4
ε	1.06	10.9
ζ	1.12	11.5
η	1.03	16.2
θ	1.27	17.3
ι	1.27	18.2
κ	1.83	24.7
λ	1.56	19.1
μ	1.92	38.0
ν	1.95	42.3
ω	.212	7.1

* Analtech Silica Gel GHLF250 μ developed with ethyl acetate:methylene chloride (1:3), detection by charring with H₂S0₄.

** Altex Ultrasphere ODS 5 μ 4.6mmx25cm developed with 85% methanol in water at 1.0 ml/minute, detection by absorbance at 254 nm.

The new agents designated LL-F28249 α , β , γ , δ , ϵ , ζ , η , θ , ι , κ , λ , μ , v and w are formed during the cultivation, under controlled conditions of Streptomyces cyaneo-griseus noncyanogenus, NRRL 15773.

This organism is maintained in the culture collection of the Medical Research Division, American Cyanamid Company, Pearl River, New York as culture number LL-F28249. A viable culture of this new microorganism has been deposited with the Patent Culture Collection Laboratory, Northern Regional Research Center, U. S. Department 10 of Agriculture, Peoria, Illinois 61604, and has been added to its permanent collection. It is freely available to the public in this depository under its accession number NRRL 15773.

For the production of these new agents the present invention is not limited to this particular organism. In fact, it is desired and intended to include the use of naturally-occurring mutants of this organism, as well as induced mutants produced from this organism by various mutagenic means known to those skilled in the art, such as exposure to nitrogen mustard, X-ray radiation, ultraviolet radiation, N'-methyl-N'-nitro-N-nitrosoguanidine, actinophages and the like. It is also desired and intended to include inter- and intraspecific genetic recombinants produced by genetic techniques known to those skilled in the art such as for example, conjugation, transduction and genetic engineering techniques.

General Fermentation Conditions

Cultivation of Streptomyces cyaneogriseus non-cyaneogenus, NRRL 15773 may be carried out in a wide variety 30 of liquid culture media. Media which are useful for the production of agents LL-F28249 α , β , γ , δ , ϵ , ζ , η , θ , ι , κ , λ , μ , v and w include an assimilable source of carbon, such as dextrin, sucrose, molasses, glycerol, etc.; an assimilable source of nitrogen such as protein, protein hydrolysate, polypeptides, amino acids, corn steep liquor, 35 etc.; and inorganic anions and cations, such as potassium, etc.;

sodium, ammonium, calcium, sulfate, carbonate, phosphate, chloride, etc. Trace elements such as boron, molybdenum, copper, etc., are supplied as impurities of other constituents of the media. Aeration in tanks and bottles is supplied by forcing sterile air through or onto the surface of the fermenting medium. Further agitation in tanks is provided by a mechanical impeller. An antifoam agent such as silicone oil may be added as needed.

Example 1

Inoculum Preparation

A typical medium used to grow the various stages of inoculum was prepared according to the following formula:

Dextrose.....	1.0%
Dextrin.....	2.0%
Yeast extract.....	0.5%
NZ amine.....	0.5%
Calcium carbonate.....	0.1%
Water.....qs.....	100%

This medium was sterilized. A 100 ml portion of this sterile medium, in a flask, was inoculated with mycelial scrapings from an agar slant of Streptomyces cyaneoegriseus noncyanogenus NRRL 15773. The medium was then agitated vigorously on a rotary shaker for 48-72 hours at 28°C providing primary inoculum. This primary inoculum was then used to inoculate one liter of the above sterile medium, which was then grown aerobically at 28°C for 48 hours providing secondary inoculum.

Example 2

Fermentation

A fermentation medium of the following formulation was prepared.

Dextrin.....	1.0%
Soya peptone.....	1.0%
Molasses.....	2.0%
Calcium carbonate.....	0.1%
Water.....qs.....	100%

This medium was sterilized and then a 30 liter portion was inoculated with one liter of secondary inoculum prepared as described in Example 1. The fermentation was conducted at 30°C, with a sterile air flow of 30 liters per minute, backpressure of 8 psig and agitation by an impeller operated at 500 rpm for 91 hours at which time the mash was harvested.

Example 3

Isolation of LL-F28249a, g andy

A total of 26 liters of whole harvest mash, prepared as described in Example 2 was mixed with 1500 g of diatomaceous earth and filtered. The mycelial cake was washed with 5 liters of water and the filtrate and wash discarded. The mycelial cake was mixed with 10 liters of methanol for one hour, then filtered and washed with 5 liters of methanol. The methanol extract and methanol wash were combined and evaporated to an aqueous residue of about 1-2 liters. This aqueous residue was mixed with twice its volume of methylene chloride and mixed for 1/2 hour. The methylene chloride phase was separated and then concentrated to a syrup giving 27 g of crude material.

This 27 g of crude material was dissolved in a mixture of methylene chloride and methanol, filtered through cotton and anhydrous sodium sulfate and then evaporated, giving 7.0 g of an oil.

A 170 g portion of silica gel was slurried in 12.5% ethyl acetate in methylene chloride and poured to form a column 2.5x58 cm. The oil was dissolved in 12.5% ethyl acetate in methylene chloride and applied to the column. The column was developed with the same solvent mixture. The mobile phase was run at 1.3 ml/minute initially and 15 minute fractions were collected. The flow rate slowed to about 0.5 ml/minute after 10 fractions, so fractions 1-10 were 20 ml decreasing to about 10 ml uniformly and fractions 11-98 were about 7 ml. At fraction 99 the flow rate was increased to give 25 ml fractions in 10 minutes. A total of 105 fractions were collected.

These fractions were tested by thin layer chromatography in ethyl acetate:methylene chloride (1:1).

Fractions 30-54 were combined and evaporated giving 1.08 g of an oil containing LL-F28249 γ .

Fractions 55-62 were combined and evaporated giving 150 mg of solid containing LL-F28249 α and β .

The 150 mg of solid containing LL-F28249 α and β was chromatographed by preparative HPLC using a reverse-phase column (Whatman C8, 2.2x50 cm) developed with 80% (v/v) methanol in water. The flow rate was about 10 ml/-10 minute and 2 minute fractions were collected.

Fractions 58-69 were combined, the methanol was evaporated, t-butanol was added and the mixture was lyophilized, giving 60 mg of pure LL-F28249 α .

Fractions 40-43 were combined, the methanol was evaporated and the residual aqueous suspension was extracted with methylene chloride which, upon evaporation, gave 10 mg of pure LL-F28249 β .

The 1.08 g of oil containing LL-F28249 γ was dissolved in 10% ethyl acetate in methylene chloride and applied to a column (2.5x50 cm) packed with silica gel. The column was developed with 10% ethyl acetate in methylene chloride, eluting at a flow rate of 2 ml/minute and collecting 12 minute fractions. Fractions 19-29 were combined and evaporated to a residue. This residue was purified by preparative reverse-phase chromatography as described for the α and β components. Fractions 55-62 were combined, the methanol was evaporated in vacuo, t-butanol was added and the mixture was lyophilized giving 60 mg of pure LL-F28249 γ .

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Example 4

Large Scale Fermentation

An inoculum of Streptomyces cyaneoegriseus non-cyanogenus, NRRL 15773 was prepared as described in Example 1, using 100 ml of primary inoculum to produce 10 liters of secondary inoculum.

Two 300 liter fermentations were conducted as described in Example 2 using 10 liters of the above secondary inoculum for each 300 liters of fermentation medium. At the end of 118 hours the mashes were harvested.

Example 5

5 Isolation of LL-F28249w

A total of 450 liters of harvest mash from the two 300 liter fermentations described in Example 4 was treated as described in the first portion of Example 3 giving crude material as a syrup.

10 This syrupy residue was washed with hexane to remove non-polar materials and the remaining 9 g of insoluble material was subjected to Sephadex LH-20 partition chromatography.

15 The chromatographic column was prepared with 9 liters of Sephadex LH-20, previously swelled in methanol, to form a column 10x110 cm. The column was equilibrated by passing about 4800 ml of mobile phase [methylene chloride:hexane:methanol (10:10:1)] through it at a flow rate of 5 ml/minute. The 9 g of insoluble material was charged onto the column in 50 ml of the mobile phase. An initial forerun of 2150 ml was obtained at a flow rate of 5 ml/minute. The flow rate was then increased to 8 ml/minute and fractions were collected every 45 minutes. Fractions 9-12 were combined and the solvents evaporated in vacuo giving 25 4.9 g of residue.

20 This residue was dissolved in a 1:1 mixture of cyclohexane and ethyl acetate and allowed to evaporate slowly at room temperature. The addition of n-hexane gave a precipitate which was collected, giving 3.1 g of solid.

25 A 3.0 g portion of this solid was further purified by precipitation from 25 ml of methylene chloride using 50 ml of n-hexane.

30 The precipitate thus obtained was redissolved in 15 ml of methylene chloride and precipitated with 25 ml of n-hexane, giving 510 mg of pure LL-F28249w.

Example 6Isolation of LL-F28249 δ , ϵ , ζ , η , θ and

Fractions 4-7 from the Sephadex LH-20 column described in Example 5 were combined and the solvents evaporated in vacuo to give 1.9 g of residue.

5 This residue was chromatographed on a 200 g silica gel column (2.5cm x 83cm) using 10% ethyl acetate in methylene chloride as the eluant. The flow rate was approximately 2 ml/minute and fractions were collected every 12 minutes.

10 Fractions 65-67 and 73-79 were combined together and the solvents were evaporated in vacuo to yield 250 mg of residue.

This 250 mg of residue was subjected to preparative reverse-phase chromatography as described in
15 Example 3 except using 75% methanol in water as the mobile phase. The flow rate was about 10 ml/minute. The first 2000 ml portion of eluate was diverted to waste then 72 fractions were collected at 2.0 minute intervals. After diverting another portion of eluate to waste (between 300-
20 400 ml) fractions were collected again but at 2.5 minute intervals.

Fractions were combined as indicated below. The combined fractions were allowed to evaporate in a fume hood overnight, then the components were extracted into
25 methylene chloride. Following evaporation of the solvent about 1 mg each of the pure components were obtained.

	<u>Fractions Combined</u>	<u>Compound</u>
	7-10	LL-F28249 δ
	19-22	LL-F28249 ϵ
30	28-31	LL-F28249 ζ
	81-83	LL-F28249 η
	86-88	LL-F28249 θ
	93-95	LL-F28249 ι

Example 7Isolation of LL-F28249 κ , λ , μ and ν

A total of 390. liters of fermentation mash, harvested from fermentations conducted as described in Example 2, was processed essentially as described in the first paragraph of Example 3, giving 120 ml of methylene chloride concentrate. This concentrate was diluted with 200 ml of hexane and chilled overnight at 4°C. The resulting precipitate was removed by filtration and discarded. The filtrate was diluted with 300 ml of hexane. The resulting precipitate (A) was collected by filtration and saved. This filtrate was evaporated to dryness and the oily residue was then dissolved in 200 ml of methylene chloride and diluted with 1700 ml of hexane. The resulting precipitate (B) was collected by filtration and saved. This filtrate was concentrated to an oily residue which was then redissolved in 50 ml of methylene chloride, 950 ml of methanol was added and this solution was stored at 4°C for 3 days. The resulting precipitate was removed by filtration and discarded. The filtrate was evaporated to dryness and the residue (C) combined with (A) and (B) and subjected to chromatography as follows: The 5.0x109cm column was slurry-packed with Woelm TSC silica gel in ethyl acetate:methylene chloride (1:9). The column was developed with the same solvent mixture at a rate of 25 ml/minute. The first 2 liters of effluent were discarded, then sixteen 400 ml fractions were collected.

Fractions 2 and 3 were combined and evaporated giving 3.9 g of oily material (D).

Fractions 4 through 7 were combined and evaporated giving 9.5 g of oily material which was dissolved in hexane and chromatographed on a 2.5x110cm column slurry-packed with 300 g of Woelm silica gel in ethyl acetate:hexane (1:4). The column was developed with the same solvent system at a rate of 4 ml/minute, collecting fractions at 7 minute intervals.

Fractions 45-54 were combined and evaporated, giving 0.3 g of material (E).

Fractions 63-135 were combined, evaporated to dryness, then redissolved in t-butanol and lyophilized giving 4.6 g of off-white solid (F).

5 LL-F28249_k and_u

Material (D) and (E) were combined and chromatographed on a 2.5x110cm column packed with 300 g of Woelm silica gel, developing with ethyl acetate:hexane (1:9). The flow rate was maintained at 4 ml/minute and fractions 10 were collected at 7 minute intervals.

Fractions 67-115 were combined and evaporated to dryness, giving 920 mg of residue (G).

This residue (G) was chromatographed by preparative HPLC using a reverse phase column (Whatman C8, 2.2x50 cm) and developing with 85% (v/v) methanol in water. The flow rate was about 10ml/minute and fractions were collected at 2.5 minute intervals.

Fractions 33-40 were combined, concentrated to remove the methanol, then extracted with methylene chloride. The residue obtained upon evaporation was dissolved in t-butanol and then lyophilized, giving 60 mg of LLF28249_k.

Fractions 52-58 were similarly processed giving a small quantity of LL-F28249_u.

25 LL-F28249_λ

A one gram portion of material (F) was chromatographed by reverse phase HPLC, as described above, except that 80% (v/v)methanol in water was used as eluent.

Fractions 61-75 were combined and processed as 30 above, giving 100 mg of LL-F28249_λ.

LL-F28249_v

A 396 g portion of material essentially the same as material (D) above, was dissolved in 500 ml of methanol and then chilled at 4° for several hours. The resulting 35 precipitate was removed by filtration, washed with cold methanol and discarded. The combined filtrate and wash

was evaporated. The residual oil was dissolved in hexane and charged on a 5x50 cm dry-packed silica gel column (Mallinkrodt SilicAR cc-7). The column was eluted with ethyl acetate:hexane (1.5:8.5) at a rate of about 50 ml/- minute.

5 Four fractions were collected.

	<u>Fraction</u>	<u>Volume(liters)</u>
	1	1
	2	4
	3	1
10	4	2

10 Fraction 3 was evaporated, giving 5.0 g of residue which was purified by preparative reverse phase HPLC (Waters C₁₈, 5x60cm). The column was initially developed with 16 liters of 80% methanol in water (v/v) at 100 ml-/minute, then with 6.4 liters of 84% methanol in water (v/v). The first liter of effluent was discarded and then fractions of 400 ml were collected.

15 Fractions 44-47 were combined and processed as described above, giving 390 mg of LLF28249v as a pale
20 yellow solid.

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Example 8Anti-nematodal activity of LL-F28249, NRRL 15773

5 This in vitro assay is designed to utilize the free living nematode Caenorhabditis elegans (C. elegans) to detect the anti-nematodal activity of fermentation broths against microorganisms from the soil. The assay procedure consists of micropipetting 50 μ l of each broth into one of 96 wells of a microculture plate and adding 10 μ l of a three to four day-old culture of C. elegans (in all stages of development) suspended in C. briggsae Maintenance Medium. The effects of the fermentation broths are observed and recorded at 48 hours after the initial mixing of broth and nematodes.

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15 LL-F28249, NRRL 15773, broth killed all the adults and markedly reduced the survival and mobility of various larval stages in both the initial and in a replicate assay.

EXAMPLE 9In vivo anthelmintic activity of LL-F28249,NRRL 15773

20 This in vivo system is designed to detect potential anthelmintic activity of all fermentation products found to have anti-nematodal action against C. elegans. Samples of LL-F28249, NRRL 15773 are mixed into feed, at concentrations of from 0.0031% to 2.0% (31 ppm to 20,000 ppm). Medicated diet containing the varying concentrations of LL-F28249, NRRL 15773 is given to gerbils infected with 400 third-stage larvae of Trichostrongylus colubriformis. The medicated feed is fed ad libitum, starting when the infection is seven days old, for three and one-half to four days, at which time the gerbils are necropsied. The intestines are removed and placed in water in an incubator at 45°C for two hours to allow the parasites to migrate from the tissue. The efficacy of each treatment is determined by counting the number of T. colubriformis recovered compared to an untreated con-

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trol. The results of these experiments, summarized in
Table XIII below, demonstrate the anthelmintic activity
of LL-F28249 as administered in feed, and when ad-
ministered as a single oral drench, and by subcutaneous
5 injection.

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TABLE XIII
**Anthelmintic activity of active ingredients from LL-F28249, NRRL 15773 culture
against Trichostrongylus colubriformis in the gerbil**

F28249				With medicated diet, Ad libitum			
Whole mash (lyophilized)	Conc. (ppm) Efficacy %	500.0 100.0	250.0 98.0	125.0 88.0	62.5 40.0		
α	Conc. (ppm) Efficacy %	20.0 100.0	0.5 100.0	0.1 97.0	0.05 31.0		
With single oral drench				With subcutaneous injection			
Whole Mash (lyophilized)	Dose (mg/kg) Efficacy %	200.0 100.0	100.0 100.0	50.0 100.0	25.0 88.0		
α	Dose (mg/kg) Efficacy %	10.0 100.0	0.5 100.0	0.1 100.0	0.05 99.0	0.025 6.0	
γ	-	-	-	0.1 78.0	0.05 15.0	0.025 10.0	
ω	-	-	-	0.1 30.0	-	-	

EXAMPLE 10The anthelmintic activity of LL-F28249a against parasitic nematodes in sheep

This experiment is designed to evaluate the activity of LL-F28249a against the economically important parasites of sheep. The sheep are experimentally inoculated with infective larvae of Haemonchus contortus, Ostertagia circumcincta and Trichostrongylus coliformis, to build up infections against which LL-F28249a will be challenged. Twenty-one days after inoculation, infection levels are determined by standard stool count nematode counting procedures to determine the number of eggs of each species per gram of feces. The sheep are assigned randomly across three replicates of treatment and control groups based upon nematode egg counts. Twenty-two days after infection the sheep are treated with LL-F28249a using the doses and routes of administration shown in Table XIV below. Seven and eight days after treatment, the sheep are sacrificed and the worms are recovered using standard anthelmintic evaluation procedures. The efficacy of each treatment against each species is determined by comparing the number of worms at the respective dosage rate against the number of worms recovered in the three untreated control animals. The results of these evaluations, summarized in Table XIV below, demonstrate the high degree of effectiveness of LL-F28249a as an anthelmintic agent.

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TABLE XIV
Anthelmintic efficacy of F28249 a
against Haemonchus, Ostertagia and Trichostrongylus in sheep

Dose mg/kg	Route of administration	Efficacy (%) against		
		Haemonchus	Ostertagia	T. colubriformis
1.0	oral	100.0	100.0	99.9
0.2	oral	100.0	100.0	99.9
	oral	100.0	95.4	99.9
0.1	oral	100.0	100.0	100.0
1.0	IM	100.0	100.0	100.0
0.2	IM	100.0	-	-
0.0	-	2683.0	881.0	16200.0
Mean number of worms recovered (range)				

IM = Intermuscular

EXAMPLE 11Efficacy of antibiotic LL-F28249 α against the parasitic insect, Melophagus ovinus, (the sheep ked) on sheep

5 This experiment is conducted concurrently on the same sheep used for the determination of anthelmintic activity as reported in Example 10. During the handling of the sheep prior to treatment, said sheep are observed for harbouring of natural infestations of M. ovinus. One half of each sheep is inspected for the indications of
10 anti-ectoparasitic activity at necropsy, seven days after treatment.

15 The left side of each sheep is slowly sheared with electric clippers and inspected for living and dead sheep keds. The degree of infestation is approximated by the numbers of pupae found in the wool during the inspection and are rated 0 through +++, indicating no pupae to many pupae. The number of keds are recorded for each sheep, without knowledge of the treatment levels to eliminate bias. Initially, the keds were scored as alive or dead,
20 but as experience was gained, some keds were scored as moribund because of abnormally-slow behavior.

25 Although there is a wide variation in the number of keds found on the sheep, the data summarized in Table XV below demonstrate that LL-F28249 α is effective against M. ovinus and that said agent possesses systemic ectoparasiticide activity. In treated animals the numbers of live keds is effectively reduced and the number of dead keds increased in the intramuscularly-treated sheep.

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TABLE XV
Efficacy of agent F28249 α against Melophagus ovinus on sheep

Dose mg/kg	Route of administration	Mean number of keds ^a		%
		Alive	Dead	
1.0	Intramuscular	1.67	1.67	78.22
	Intramuscular	1.0	4.33	86.96
0.2	Oral	7.67	0.0	0.0
			65.0	65.0
1.0	Oral	2.67	3.0	0.0
0.2	Oral	22.0	1.67	-
0.1	Oral	.67	.67	-
Control	None	7.67		

^a Three sheep per dose Mean number in control - mean number in treated
^b Efficacy % = 100 X $\frac{\text{Mean number in treated}}{\text{Mean number in control}}$

EXAMPLE 12Insecticidal activity of the compounds of the invention

5 The insecticidal activity of the compounds of the present invention against a variety of insects at various concentrations of active ingredient in acetone-water solutions is determined by the following insecticidal test examples. The results of these tests are summarized in Table XVI.

10 A) Heliothis virescens, egg, tobacco budworm. A young cotton leaf about 7-8 cm long is dipped and agitated in a test suspension for three seconds. Eggs are collected on cheesecloth that is cut into 10-20 mm squares containing about 50-100 eggs (6-30 hours old). A square of cheesecloth with eggs also is dipped in the test suspension and placed on the treated leaf. The combination is placed in the hood to dry. Following this, the combination is placed in an 8 ounce Dixie cup #2168-ST (240 mL, 6 cm tall, top diameter 9.5 cm, bottom diameter 8 cm) containing a 5 cm length of damp dental wick. A clear plastic lid is put on the top of the cup, and the treatments held for three (3) days before mortality counts are made.

15 B) Aphis fabae, mixed instars, bean aphids. Pots containing single masturtium plant (Tropaeolum sp), about 5 cm tall, are infested with about 100 aphids one day before the test. In a hood, each plant is sprayed with the test suspension for 2 revolutions of a 4 rpm turntable using a #154 DeVilluss atomizer. The pots are set on their side on white enamel trays and held for two (2) days. After that time, mortality estimates of the aphids are made.

20 C) Emoasca abrupta, adult, western potato leafhopper. A Sieva lima bean leaf about 5 cm long is dipped and agitated in the test suspension for three (3) seconds and then placed in a hood to dry. The leaf is placed in a 100 x 10 mm petri dish containing a moist filter paper on the

bottom of the dish. Ten, adult leafhoppers are added to each dish, and the treatments are kept for three (3) days after which time mortality counts are made.

D) Trichoplusia ni, Third-instar larvae, cab-

5 bage looper.

The leaves of a Sieva lima bean plant expanded to 7-8 cm in length are dipped and agitated in a test suspension for three (3) seconds and then placed in a hood to dry. A leaf is then excised and placed in a 100 x 10 mm petri dish containing a damp filter paper on the bottom and ten third-instar larvae are placed therein. The dish is maintained for three (3) days before observations are made of mortality and reduced feeding.

E) Spodoptera eridania, third-instar larvae,

15 southern armyworm.

The leaves of a Sieva lima bean plant expanded to 7-8cm in length are dipped and agitated in the test suspension for three (3) seconds and placed in a hood to dry. A leaf is then excised and placed in a 100 x 10 mm petri dish containing a damp filter paper on the bottom and ten (10) third-instar larvae are added. The dish is maintained for five (5) days before observations are made of mortality, reduced feeding or any interference with normal moulting.

F) Heliothis virescens, third-instar larvae,
tobacco budworm.

Cotton cotyledons are dipped in the test suspension and placed in a hood to dry. The cotyledon is cut into 4 sections, and each section is placed in a 30 ml plastic medicine cup containing a 5-7 mm piece of moist dental wick. One third-instar larvae are added to each cup and a cardboard lid placed on the cup. Treatments are maintained for three (3) days before mortality counts and estimates of reduction in feeding are made.

G) Musca domestica, house fly.

The desired concentration of the test compound is added to the standard CSMA alfalfa-bran larval medium. House

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flies' eggs, 0-4 hours of age, are added to the treated medium. The treated medium is maintained and observations on egg hatch, larval growth and adult emergence are made.

5 H) Tribolium confusum, confused flour beetle. Confused flour beetles (Tribolium confusum) are obtained from laboratory colonies reared on a whole wheat and white flour mixture. For this test, white flour is treated with an acetone solution of the test material using 1 ml of solution per 5 grams of flour in a 30 ml wide-mouth jar. The acetone is evaporated off in a hood overnight. The contents are stirred with a spatula to break up lumps formed by the test solution. The jar is then placed on a VORTES-GENIE® vibrating mixer to thoroughly mix the test materials throughout the diet. Ten adult confused flour beetles are placed in each jar and the jar loosely capped. After five (5) days to allow oviposition, the beetles are removed and notations made of any mortality. At two (2) and four (4) weeks after initial infestation, observations are made of the number and size of trails produced by the developing larvae throughout the treated flour. Such observations give an indication of delayed growth, kill of eggs or larvae or any other interference in the normal growth pattern. After about nine (9) weeks at 27°C, the adult beetles emerge and the final observations are made by passing the contents of each jar through a 50-mesh screen sieve. These observations include the number of adults, pupae and larvae, as well as examination of the debris which did not pass through the screen in order to determine if there are any dead eggs or neonates.

20 I) Tetranychus urticae (P-resistant strain), 2-spotted spider mite.
25 Sieva lima bean plants with primary leaves expanded to 7-8 cm are selected and cut back to one plant per pot. A small piece is cut from a leaf taken from the main colony
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and placed on each leaf of the test plants. This is done about two (2) hours before treatment to allow the mites to move over to the test plant and to lay eggs. The size of the cut piece is varied to obtain about 100 mites per leaf. At the time of the treatment, the piece of leaf used to transfer the mites is removed and discarded. The mite-infested plants are dipped and agitated in the test formulation for three (3) seconds and set in the hood to dry. Plants are kept for two (2) days before estimates of adult kill are made by using the first leaf. The second leaf is kept on the plant for another five (5) days before observations are made of the kill of eggs and/or newly emerged nymphs.

5 J) Southern armyworm (Spodoptera eridania),
10 third-instar, cut-stem systemic test.
15 The compound is formulated as an emulsion containing 0.1 gm of the test material, 0.1 gm of a polyethoxylated vegetable oil in 0.4 g water, 10 mL of acetone and 90 mL of water. This is diluted ten-fold with water to give the
20 100 ppm emulsion for the test. Sieva lima bean plants with just the primary leaves expanded are used in this test. These leaves are cut off at least 2.5 cm above the soil level to avoid contamination with soil bacteria which may cause decay of the stem during the test. The
25 cut stems are placed in the test emulsion. After three (3) days of uptake, a leaf is excised and placed in a 100 x 10 mm petri dish containing a moist filter paper on the bottom and ten third-instar larvae. Mortality counts and estimates of reduced feeding are made after three (3)
30 days.

35 K) Thrips palmi, thrips.
Heavily infested leaves of cotton seedlings are sprayed under field conditions at the desired concentrations. The number of thrips are counted before and after spraying. Percent control is based on these counts.
L) Tetranychus urticae (P-resistant strain), two spotted spider mite.

The compound is formulated as an emulsion containing 0.1 gm of the test material, 0.1 gm of a polyethoxylated vegetable oil in 0.4 g water, 10 mL of acetone and 90 mL of water. This is diluted ten-fold with water to give the 100 ppm emulsion for the test. Sieva lima bean plants with just the primary leaves expanded are used in this test. They are cut off at least 2.5 cm above the soil level to avoid contamination with soil bacteria which may cause decay of the stem during the test. The cut stems are placed in the test emulsions. Each leaf is infested with approximately 100 adult mites and maintained for three (3) days at which time mortality counts are made.

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TABLE XVI
Insecticidal and Miticidal Activity of F-28,249 α and F-28,249 γ

Compound	Concn. in ppm	Percent Mortality										Plant Systemic Activity
		Cabbage loopers	Southern army- worms	Tobacco bud- worms	Bean eggs	Bean aphid	Western Confused flour beetle larvae and/or pupae	Potato leaf hoppers	House fly larvae	Thrips	Mites	
F-28,249 α 1000	100	100	100	-	-	55*	100	100	93	-	-	-
F-28,249 α 300	100*	-	100	100	-	50	-	100	89	100	-	-
F-28,249 α 100	-	60*	100*	100	100	-	97	-	-	100	60	100
F-28,249 γ 1000	-	40	50*	100	100	20	-	-	-	100	-	-
F-28,249 γ 100	-	0	0	0	100	0	-	-	-	90	-	-

* Feeding deterrent (anti-feeding properties)

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WHAT IS CLAIMED IS:

1. The compound designated LL-F28249a wherein
the compound has:
- a) a molecular weight of 612 (FAB-MS);
 - b) a molecular formula, C₃₆H₅₂O₈;
 - 5 c) a specific optical rotation, [α]_D²⁶ = +133±3° (C 0.3,
acetone);
 - d) a characteristic ultraviolet absorption spectrum as
shown in Figure I of the attached drawings;
 - e) a characteristic infrared absorption spectrum as shown
10 in Figure II of the attached drawings;
 - f) a characteristic proton nuclear magnetic resonance
spectrum as shown in Figure III of the attached draw-
ings;
 - 15 g) a characteristic carbon-13 nuclear magnetic resonance
spectrum as shown in Figure IV of the attached draw-
ings with significant peaks at, 173.4; 142.8; 139.4;
137.7; 137.3; 137.2; 130.6; 123.3; 120.3; 118.0; 99.7;
80.2; 79.3; 76.7; 69.3; 68.5; 68.4; 67.8; 67.7; 48.4;
45.7; 41.1; 40.7; 36.1; 36.0; 35.9; 34.7; 26.8; 22.8;
20 22.2; 19.9; 15.5; 13.9; 11.0; and
 - 25 h) a characteristic electron impact mass spectrum as
shown in Figure V of the attached drawings with mea-
sured m/z values and proposed elemental compositions
as indicated below obtained by high resolution mass
measurements,
- | | | | |
|-------------|--|----------|---|
| 612.3705 | C ₃₆ H ₅₂ O ₈ | 354.2181 | C ₂₃ H ₃₀ O ₃ |
| 594.3543 | C ₃₆ H ₅₀ O ₇ | 314.1877 | C ₂₀ H ₂₆ O ₃ |
| 576.3472 | C ₃₆ H ₄₈ O ₆ | 278.1144 | C ₁₅ H ₁₈ O ₅ |
| 484.3211 | C ₃₀ H ₄₄ O ₅ | 265.1786 | C ₁₆ H ₂₅ O ₃ |
| 30 482.2648 | C ₂₉ H ₃₈ O ₆ | 248.1405 | C ₁₅ H ₂₀ O ₃ |
| 466.3097 | C ₃₀ H ₄₂ O ₄ | 247.1705 | C ₁₆ H ₂₃ O ₂ |
| 448.2987 | C ₃₀ H ₄₀ O ₃ | 237.1838 | C ₁₅ H ₂₅ O ₂ |
| 442.2375 | C ₂₆ H ₃₄ O ₆ | 219.1740 | C ₁₅ H ₂₃ O |
| 425.2327 | C ₂₆ H ₃₃ O ₅ | 151.0753 | C ₉ H ₁₁ O ₂ . |

2. The compound designated LL-F28249₈ wherein
the compound has:
- a) a molecular weight of 584 (FAB-MS);
 - b) a molecular formula, C₃₄H₄₈O₈;
 - c) specific optical rotation: [α]_D²⁶ = +125° (C 0.30, acetone).
 - 5 d) a characteristic ultraviolet absorption spectrum as shown in Figure VI of the attached drawings;
 - e) a characteristic infrared absorption spectrum as shown in Figure VII of the attached drawings;
 - 10 f) a characteristic proton nuclear magnetic resonance spectrum as shown in Figure VIII of the attached drawings;
 - g) a characteristic carbon-13 nuclear magnetic resonance spectrum as shown in Figure XXXVIII of the attached drawings, with significant peaks at 173.3; 142.6; 139.5; 137.7; 137.3; 133.9; 123.8; 123.4; 120.3; 120.2; 118.0; 99.7; 80.2; 79.4; 76.7; 69.2; 68.6; 68.3; 67.8; 67.7; 48.4; 45.7; 41.0; 40.8; 36.1; 35.9; 34.7; 22.3; 19.8; 15.5; 13.8; 13.1; 10.8; and
 - 15 h) a characteristic electron impact mass spectrum as shown in Figure IX of the attached drawings with measured m/z values and proposed elemental compositions as indicated below obtained by high resolution mass measurements,
 - 20 584.3388 C₃₄H₄₈O₈ 314.1858 C₂₀H₂₆O₃
 - 566.3306 C₃₄H₄₆O₇ 278.1168 C₁₅H₁₈O₅
 - 456.2864 C₂₈H₄₀O₅ 237.1491 C₁₄H₂₁O₃
 - 442.2391 C₂₆H₃₄O₆ 219.1380 C₁₄H₁₉O₂
 - 438.2780 C₂₈H₃₈O₄ 209.1534 C₁₃H₂₁O₂
 - 30 425.2331 C₂₆H₃₃O₅ 191.1418 C₁₃H₁₉O
 - 354.2187 C₂₃H₃₀O₃ 151.0750 C₉H₁₁O₂.
3. The compound designated LL-F28249_Y wherein
the compound has:
- a) a molecular weight of 598 (FAB-MS);
 - b) a molecular formula, C₃₅H₅₀O₈;
 - c) a specific optical rotation: [α]_D²⁶ = +150±4° (C 0.3, acetone);

- d) a characteristic ultraviolet absorption spectrum as shown in Figure X of the attached drawings;
- e) a characteristic infrared absorption spectrum as shown in Figure XI of the attached drawings;
- f) a characteristic proton nuclear magnetic resonance spectrum as shown in Figure XII of the attached drawings;
- 5 g) a characteristic carbon-13 nuclear magnetic resonance spectrum as shown in Figure XIII of the attached drawings with significant peaks at, 173.6; 142.4; 139.9; 10 137.3; 136.0; 134.0; 123.8; 123.6; 120.4; 119.6; 118.5; 99.8; 80.5; 77.8; 77.0; 76.8; 69.3; 68.6; 68.3; 67.9; 57.7; 48.5; 45.8; 41.2; 40.8; 36.2; 36.1; 36.0; 34.8; 22.3; 19.9; 15.5; 13.8; 13.1; 10.8; and 15 h) a characteristic electron impact mass spectrum as shown in Figure XIV of the attached drawings with measured m/z values and proposed elemental compositions as indicated below obtained by high resolution mass measurements,

598.3543	C ₃₅ H ₅₀ O ₈	354.2199	C ₂₃ H ₃₀ O ₃
20 580.3422	C ₃₅ H ₄₈ O ₇	314.1875	C ₂₀ H ₂₆ O ₃
562.3292	C ₃₅ H ₄₆ O ₆	292.1307	C ₁₆ H ₂₀ O ₅
496.2824	C ₃₀ H ₄₀ O ₆	288.2175	C ₁₉ H ₂₈ O ₂
484.2440	C ₂₈ H ₃₆ O ₇	248.1397	C ₁₅ H ₂₀ O ₃
478.2687	C ₃₀ H ₃₈ O ₅	237.1490	C ₁₄ H ₂₁ O ₃
25 456.2576	C ₂₇ H ₃₆ O ₆	219.1282	C ₁₄ H ₁₉ O ₂
438.2772	C ₂₈ H ₃₈ O ₄	209.1544	C ₁₃ H ₂₁ O ₂
425.2341	C ₂₆ H ₃₃ O ₅	191.1435	C ₁₃ H ₁₉ O
420.2651	C ₂₈ H ₃₆ O ₃	151.0759	C ₉ H ₁₁ O ₂ .

4. The compound designated LL-F28249_w wherein
30 the compound has:
- a) a molecular weight of 806 (FAB-MS);
- b) a molecular formula, C₄₅H₇₄O₁₂;
- c) a specific optical rotation: [α]_D²⁶ = -49^{±40} (C 0.35, methanol);
- 35 d) a characteristic ultraviolet absorption spectrum as shown in Figure XV of the attached drawings;

- e) a characteristic infrared absorption spectrum as shown in Figure XVI of the attached drawings;
 - f) a characteristic proton nuclear magnetic resonance spectrum as shown in Figure XVII of the attached drawings;
 - 5 g) a characteristic carbon-13 nuclear magnetic resonance spectrum as shown in Figure XVIII of the attached drawings with significant peaks at, 220.7; 219.6; 165.2; 148.7; 133.1; 132.3; 130.2; 122.3; 100.0; 82.9; 75.9; 73.0; 72.7; 72.6; 72.1; 69.0; 67.3; 63.6; 51.4; 46.2; 45.7; 42.2; 40.4; 38.3; 37.6; 36.1; 34.8; 33.5; 30.1; 26.6; 25.4; 24.5; 23.0; 21.1; 17.9; 14.3; 14.2; 12.1; 11.5; 10.9; 8.7; 8.3; 5.7; and
 - 10 h) a characteristic electron impact mass spectrum as shown in Figure XIX of the attached drawings with measured m/z values and proposed elemental compositions as indicated below obtained by high resolution mass measurements,
 - 15

462.3350	C ₂₈ H ₄₆ O ₅	253.1797	C ₁₅ H ₂₅ O ₃
444.3237	C ₂₈ H ₄₄ O ₄	235.1697	C ₁₅ H ₂₃ O ₂
20 425.2534	C ₂₃ H ₃₇ O ₇	224.1754	C ₁₄ H ₂₄ O ₂
407.2439	C ₂₃ H ₃₅ O ₆	209.1530	C ₁₃ H ₂₁ O ₂
406.3046	C ₂₅ H ₄₂ O ₄	207.1744	C ₁₄ H ₂₃ O
387.2895	C ₂₅ H ₃₉ O ₃	184.1458	C ₁₁ H ₂₀ O ₂
337.2010	C ₁₉ H ₂₉ O ₅	179.1048	C ₁₁ H ₁₅ O ₂
25 297.2031	C ₁₇ H ₂₉ O ₄	173.1205	C ₉ H ₁₇ O ₃
279.1944	C ₁₇ H ₂₇ O ₃	167.1051	C ₁₀ H ₁₅ O ₂
261.1851	C ₁₇ H ₂₅ O ₂	155.1069	C ₉ H ₁₅ O ₂ .
5. The compound designated LL-F28249 & wherein

- the compound has:
- 30 a) a molecular weight of 616 (EI-MS);
 - b) a molecular formula, C₃₅H₅₂O₉;
 - c) a HPLC retention volume of 14.0 ml;
 - d) a characteristic ultraviolet absorption spectrum shown in Figure XX of the attached drawings;
 - 35 e) a characteristic proton nuclear magnetic resonance spectrum as shown in Figure XXI of the attached draw-

- f) a characteristic electron impact mass spectrum as shown in Figure XXII of the attached drawings.

6. The compound designated LL-F28249 ϵ wherein the compound has:
- a) a molecular weight of 598 (EI-MS);
 - 5 b) molecular formula, C₃₅H₅₀O₈;
 - c) a HPLC retention volume of 14.8 ml;
 - d) a characteristic ultraviolet absorption spectrum as shown in Figure XXIII of the attached drawings;
 - e) a characteristic proton nuclear magnetic resonance spectrum as shown in Figure XXIV of the attached drawings; and
 - 10 f) a characteristic electron impact mass spectrum as shown in Figure XXV of the attached drawings.

7. The compound designated LL-F28249 ζ wherein the compound has:
- a) a molecular weight of 598 (EI-MS);
 - b) a molecular formula, C₃₅H₅₀O₈;
 - c) a HPLC retention volume of 16.0 ml;
 - d) a characteristic ultraviolet absorption spectrum as shown in Figure XXVI of the attached drawings;
 - 20 e) a characteristic proton nuclear magnetic resonance spectrum as shown in Figure XXVII of the attached drawings; and
 - f) a characteristic electron impact mass spectrum as shown in Figure XXVIII of the attached drawings.

- 25 8. The compound designated LL-F28249 η wherein the compound has:
- a) a molecular weight of 612 (EI-MS);
 - b) a molecular formula, C₃₆H₅₂O₈;
 - 30 c) a HPLC retention value of 23.5 ml;
 - d) a characteristic ultraviolet absorption spectrum as shown in Figure XXIX of the attached drawings;
 - e) a characteristic proton nuclear magnetic resonance spectrum as shown in Figure XXX of the attached drawings; and
 - 35 f) a characteristic electron impact mass spectrum as

shown in Figure XXXI of the attached drawings.

9. The compound designated LL-F28249₀ wherein the compound has:

- a) a molecular weight of 626 (EI-MS);
- b) a molecular formula, C₃₇H₅₄O₈;
- 5 c) a HPLC retention value of 24.5 ml;
- d) a characteristic ultraviolet absorption spectrum as shown in Figure XXXII of the attached drawings;
- e) a characteristic proton nuclear magnetic resonance spectrum as shown in Figure XXXIII of the attached
- 10 drawings; and
- f) a characteristic electron impact mass spectrum as shown in Figure XXXIV of the attached drawings.

10. The compound designated LL-F28249₁ wherein the compound has:

- 15 a) a molecular weight of 626 (EI-MS);
- b) a molecular formula, C₃₇H₅₄O₈;
- c) a HPLC retention value of 26.0 ml;
- d) a characteristic ultraviolet absorption spectrum as shown in Figure XXXV of the attached drawings;
- 20 e) a characteristic proton nuclear magnetic resonance spectrum as shown in Figure XXXVI of the attached drawings; and
- f) a characteristic electron impact mass spectrum as shown in Figure XXXVII of the attached drawings.

25 11. The compound designated LL-F28249_k wherein the compound has:

- a) molecular weight 584 (EI-MS);
- b) molecular formula: C₃₅H₅₂O₇;
- 30 c) Specific optical rotation: [α]_D²⁶=+189°- (C 0.165 acetone);
- d) Ultraviolet absorption spectrum: as shown in Figure XXXIX UV _{CH₃OH} MAX_{241nm} (E20,400);
- e) Infrared absorption spectrum: as shown in Figure XL (KBr disc);
- 35 f) Electron impact mass spectrum: as shown in Figure XLI;

- g) Proton nuclear magnetic resonance spectrum (CDCl₃); as shown in Figure XLII; and
h) Carbon-13 nuclear magnetic resonance spectrum (CDCl₃); as shown in Figure XLIII and described in Table IX.

12. The compound designated LL-F28249λ wherein

- 5 the compound has:
- a) molecular weight: 626 (FAB-MS);
 - b) Molecular formula: C₃₇H₅₄O₈;
 - c) specific optical rotation: [α]_D²⁶+145°(C, 0.23 acetone);
 - d) Ultraviolet absorption spectrum: as shown
10 in Figure XLIV UV _{CH₃OH}^{MAX}=244nm (E30,000);
 - e) Infrared absorption spectrum: as shown in Figure XLV (KBr disc);
 - f) Electron impact mass spectrum: as shown in Figure
15 XLVI;
 - g) Proton nuclear magnetic resonance spectrum (CDCl₃); as shown in Figure XLVII; and
 - h) Carbon-13 nuclear magnetic resonance spectrum (CDCl₃); as shown in Figure XLVIII and described in Table X.

13. The compound designated LL-F28249μ wherein

- 20 the compound has:
- a) molecular weight: 612 (EI-MS);
 - b) molecular formula: C₃₇H₅₆O₇;
 - c) Ultraviolet absorption spectrum: as shown
25 in Figure XLIX UV _{CH₃OH}^{MAX}=241nm (E16,800);
 - d) Infrared absorption spectrum: as shown in Figure L (KBr disc);
 - e) Electron impact mass spectrum: as shown in Figure LI;
 - f) Proton nuclear magnetic resonance spectrum (CDCl₃); as shown in Figure LII.

30 14. The compound designated LL-F28249ν wherein
the compound has:

- a) molecular weight: 592 (EI-MS);
- b) molecular formula: C₃₆H₄₈O₇;
- c) specific optical rotation: [α]_D²⁶+131°(C .325, acetone);
- 35 d) Ultraviolet absorption spectrum: as shown in Figure

- LIII UV CH_3OH _{MAX} = 256 (E 20,500); 358 (E 8,830);
- e) Infrared absorption spectrum: as shown in Figure LIV (KBr disc);
- f) Electron impact mass spectrum: as shown in Figure LV;
- 5 g) Proton nuclear magnetic resonance spectrum (CDCl_3); as shown in Figure LVI; and
- h) Carbon-13 nuclear magnetic resonance spectrum (CDCl_3); as shown in Figure LVII, and described in Table XI.
15. A process for producing agents LL-F28249 α , LL-F28249 β , LL-F28249 γ , LL-F28249 δ , LL-F28249 ϵ , LL-F28249 ζ , LL-F28249 η , LL-F28249 θ , LL-F28249 ι , LL-F28249 κ , LL-F28249 λ , LL-F28249 μ , LL-F28249 ν , and LL-F28249 ω which comprises: aerobically fermenting the organism Streptomyces cyaneogriseus noncyanogenus, NRRL 15773 or an LL-F28249 α , β , γ , δ , ϵ , ζ , η , θ , ι , κ , λ , μ , ν and ω producing mutant thereof, in a liquid medium containing assimilable sources of carbon, nitrogen and inorganic anions and cations, until a substantial amount of LL-F28249 α , β , γ , δ , ϵ , ζ , η , θ , ι , κ , λ , μ , ν and ω are produced in said medium; and then recovering the 20 agents therefrom.
16. A process for producing agents LL-F28249 α , LL-F28249 β , LL-F28249 γ , LL-F28249 δ , LL-F28249 ϵ , LL-F28249 ζ , LL-F28249 η , LL-F28249 θ , LL-F28249 ι , LL-F28249 κ , LL-F28249 λ , LL-F28249 μ , LL-F28249 ν , and LL-F28249 ω which comprises: aerobically fermenting a liquid medium containing assimilable sources of carbon, nitrogen and inorganic anions and cations, which medium has been inoculated with a viable culture of the organism Streptomyces cyaneogriseus noncyanogenus, NRRL 15773 or an LL-F28249 α , β , γ , δ , ϵ , ζ , η , θ , ι , κ , λ , μ , ν and ω producing mutant thereof; maintaining said fermentation culture with sterile aeration and agitation at a temperature of 24°-32°C for a period of 25-200 hours; harvesting the mash; and extracting the 30 agents.
- 35 17. A biologically pure culture of the micro-organism Streptomyces cyaneogriseus noncyanogenus, NRRL

15773, said culture being capable of producing agents LL-F28249 α , LL-F28249 β , LL-F28249 γ , LL-F28249 δ , LL-F28249 ϵ , LL-F28249 ζ , LL-F28249 η , LL-F28249 θ , LL-F28249 ι , LL-F28249 κ , LL-F28249 λ , LL-F28249 μ , LL-F28249 ν , and LL-F28249 ω in recoverable quantities upon fermentation in an aqueous nutrient medium containing assimilable sources of carbon, nitrogen and in-organic anions and cations.

18. The biologically pure culture of the micro-organism Streptomyces cyaneogriseus noncyanogenus, according to Claim 13, wherein said microorganism has spontaneously mutated, such that the microorganism is genetically altered but still retains the ability to synthesize agents LL-F28249 α , LL-F28249 β , LL-F28249 γ , LL-F28249 δ , LL-F28249 ϵ , LL-F28249 ζ , LL-F28249 η , LL-F28249 θ , LL-F28249 ι , LL-F28249 κ , LL-F28249 λ , LL-F28249 μ , LL-F28249 ν , and LL-F28249 ω .

19. The biologically pure culture of the microorganism Streptomyces cyaneogriseus noncyanogenus, according to Claim 13, wherein said microorganism has been subjected to mutagenic means such that the microorganism is genetically altered but still retains the ability to synthesize agents LL-F28249 α , LL-F28249 β , LL-F28249 γ , LL-F28249 δ , LL-F28249 ϵ , LL-F28249 ζ , LL-F28249 η , LL-F28249 θ , LL-F28249 ι , LL-F28249 κ , LL-F28249 λ , LL-F28249 μ , LL-F28249 ν , and LL-F28249 ω .

20. The compound according to Claim 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13 or 14 additionally comprising: the pharmaceutically and pharmacologically-acceptable salts thereof.

21. A method for the prevention, treatment or control of helmintic, arthropod ectoparasitic or acaridal infections in warm-blooded animals, said method comprising: orally, parenterally or topically administering to an animal infected with helminths, arthropod ectoparasites or acarides, a prophylactically, therapeutically or pharmaceutically-effective amount of the fermentation broth or whole mash of micro-organism Streptomyces cyaneogriseus noncyanogenus, having deposit accession number NRRL 15773.

22. A method for the prevention, treatment or control of helmintic, arthropod ectoparasitic or acaridal infections

in warm-blooded animals, said method comprising: orally, parenterally or topically administering to an animal infected with helminths, arthropod ectoparasites or acarides, a prophylactically, therapeutically or pharmaceutically-effective amount of the fermentation broth or whole mash of microorganism Streptomyces sp. LL-F28249, having deposit accession number NRRL 15773, containing agents designated LL-F28249 α , LL-F28249 β , LL-F28249 γ , LL-F28249 δ , LL-F28249 ϵ , LL-F28249 ζ , LL-F28249 η , LL-F28249 θ , LL-F28249 ι , LL-F28249 κ , LL-F28249 λ , LL-F28249 μ , LL-F28249 ν , and LL-F28249 ω ; or pharmaceutically and pharmacologically-acceptable salts thereof.

10 23. A method for the treatment of helminthic infections according to Claims 21 or 22, wherein about 0.1 mg per kg to 200 mg per kg is administered to an animal infected with helminths.

15 24. A method for the control of plant nematodes, said method comprising: applying to the foliage of plants, the soil in which they are grown, or into the trunks thereof, a nematocidally-effective amount of the fermentation broth or whole mash of microorganism Streptomyces cyaneogriseus non-cyanogenus, having deposit accession number NRRL 15773.

20 25. A method for the control of plant nematodes, said method comprising: applying to the foliage of plants, the soil in which they are grown, or into the trunks thereof, a nematocidally-effective amount of the fermentation broth or whole mash of microorganism Streptomyces cyaneogriseus non-cyanogenus, having deposit accession number NRRL 15773, containing agents designated LL-F28249 α , LL-F28249 β , LL-F28249 γ , LL-F28249 δ , LL-F28249 ϵ , LL-F28249 ζ , LL-F28249 η , LL-F28249 θ , LL-F28249 ι , LL-F28249 κ , LL-F28249 λ , LL-F28249 μ , LL-F28249 ν , and LL-F28249 ω ; or the pharmaceutically and pharmacologically-acceptable salts thereof.

25 26. A method according to Claims 24 or 25, wherein about 0.1 to 1.4 kg per hectare is applied to thereof.

30 27. An animal feed composition for the prevention, treatment or control of helmintic, arthropod ectoparasitic or acaridal infections in meat-producing animals, said animal feed composition comprising: an edible solid carrier; and a prophylactically, therapeutically or pharmaceutically-

effective amount of the fermentation broth or whole mash of microorganism Streptomyces cyaneogriseus noncyanogenus, having deposit accession number NRRL 15773.

28. An animal feed premix composition for the prevention, treatment or control of helminthic, arthropod ectoparasitic or acaridal infections in meat-producing animals, said animal feed premix composition comprising: an edible carrier; and a prophylactically, therapeutically or pharmaceutically-effective amount of the fermentation broth or whole mash of microorganism Streptomyces cyanogriseus noncyanogenus, having deposit accession number NRRL 15773, containing agents designated LL-F28249 α , LL-F28249 β , LLF28249 γ , LL-F28249 δ , LL-F28249 ϵ , LL-F28249 ζ , LL-F28249 η , LL-F28249 θ , L1-F28249 ι , LL-F28249 κ , LL-F28249 λ , LL-F28249 μ , LL-F28249 ν and LL-F28249 ω ; or the pharmaceutically and pharmacologically acceptable salts thereof.

29. A composition according to Claims 27 or 28, wherein said effective amount is about 0.00001% to 5%, by weight, of said composition.

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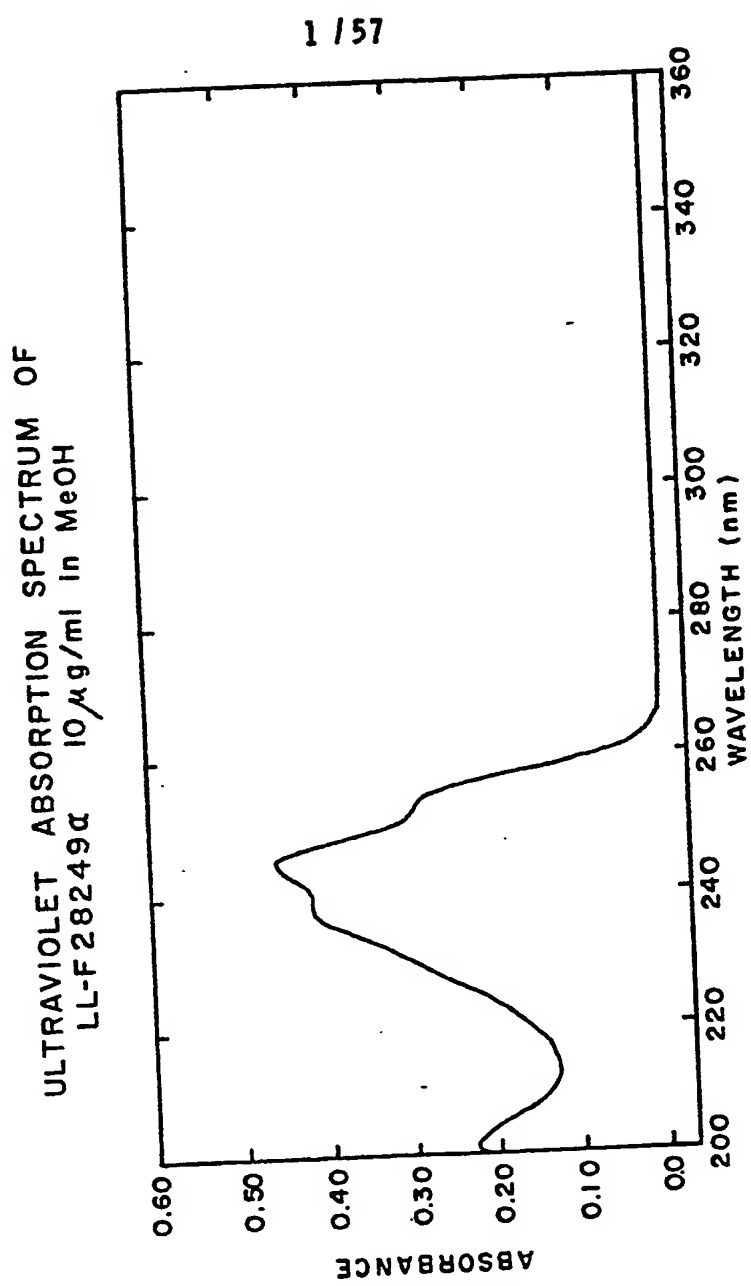


FIGURE 1

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INFRARED ABSORPTION SPECTRUM
OF LL-F28249a (KBr disc)

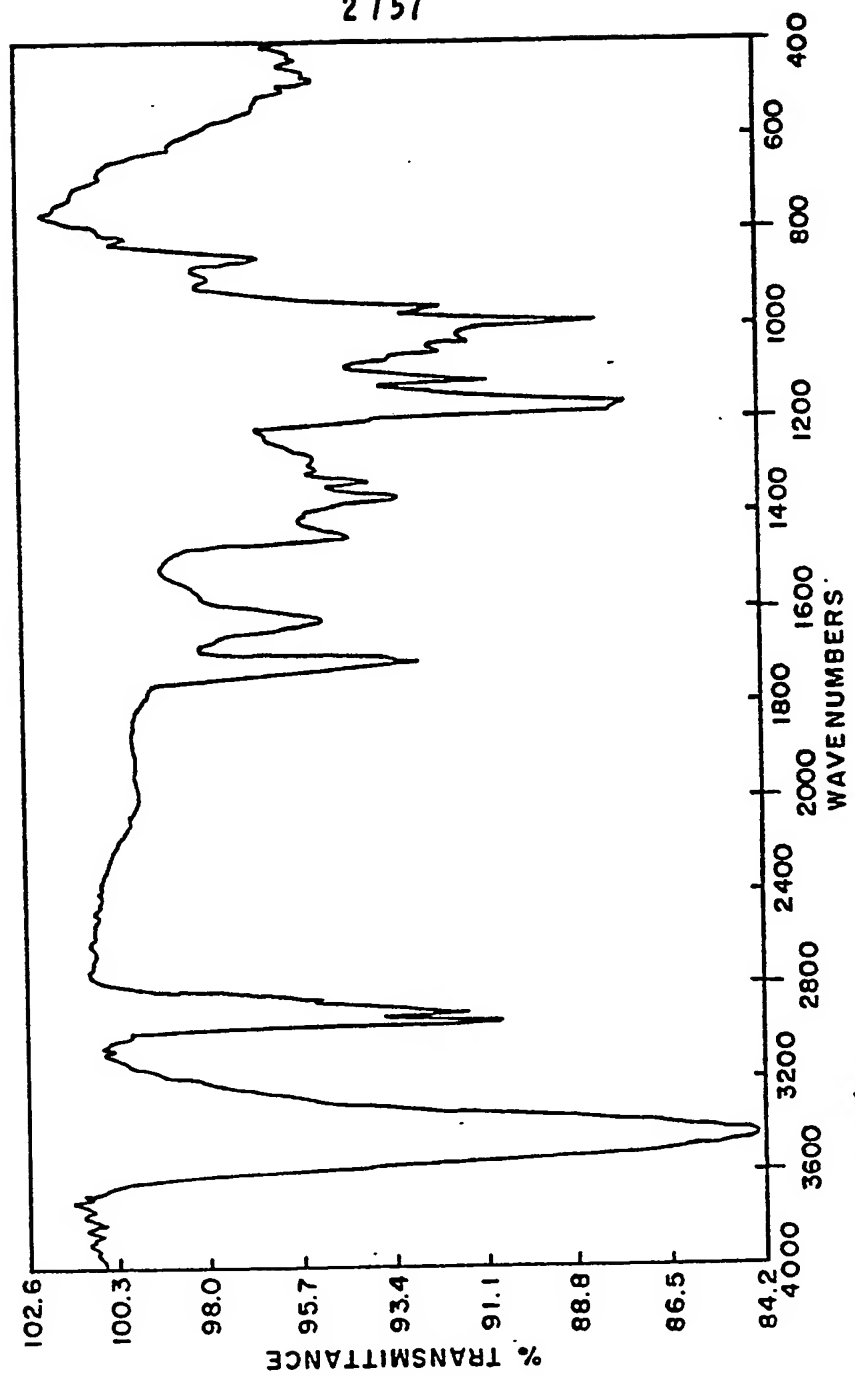


FIGURE 2

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PROTON NUCLEAR MAGNETIC RESONANCE SPECTRUM
OF LL-F28249 α IN CDCl₃ SOLUTION

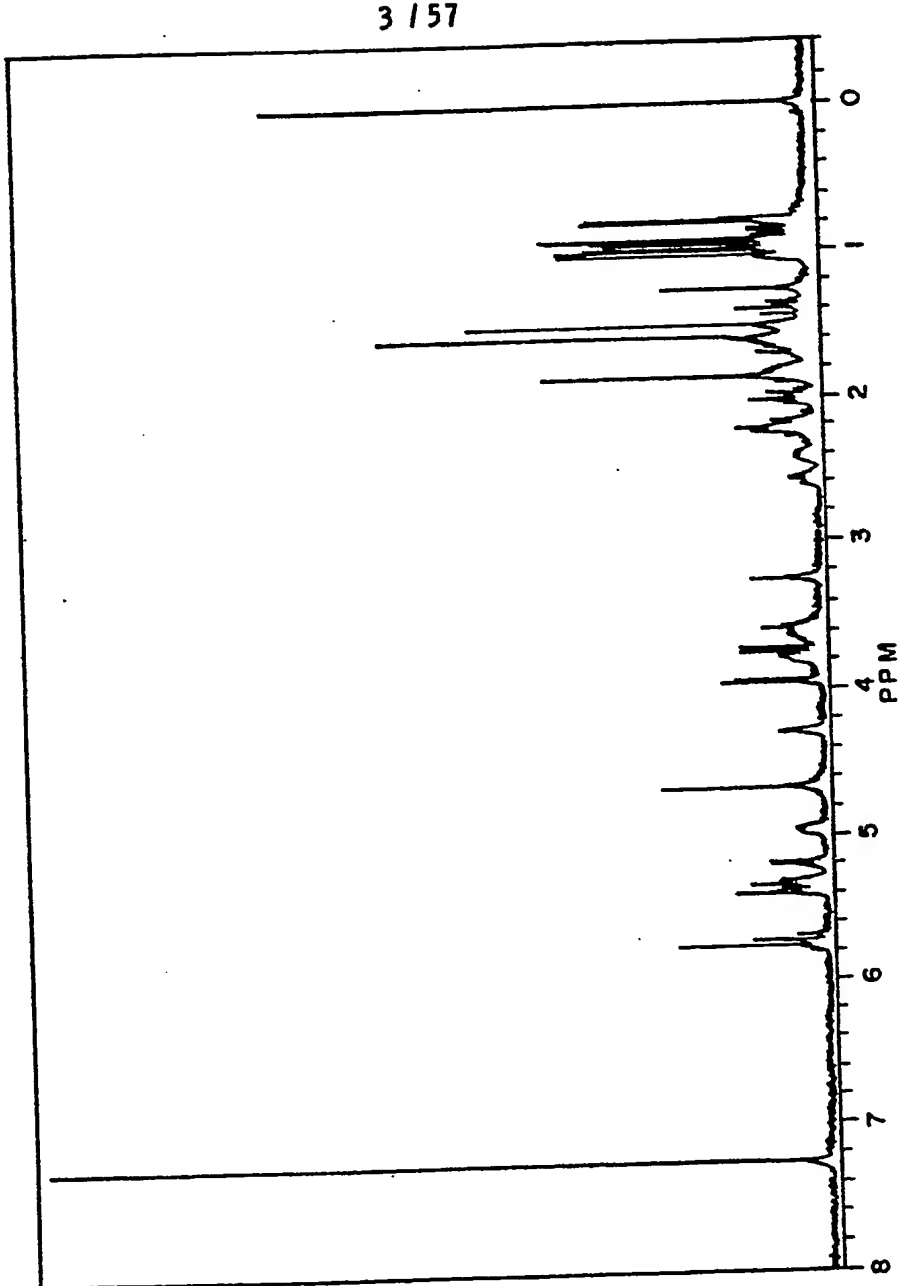


FIGURE 3

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CARBON-13 NUCLEAR MAGNETIC RESONANCE SPECTRUM
OF LL-F28249d IN CDCl₃ SOLUTION

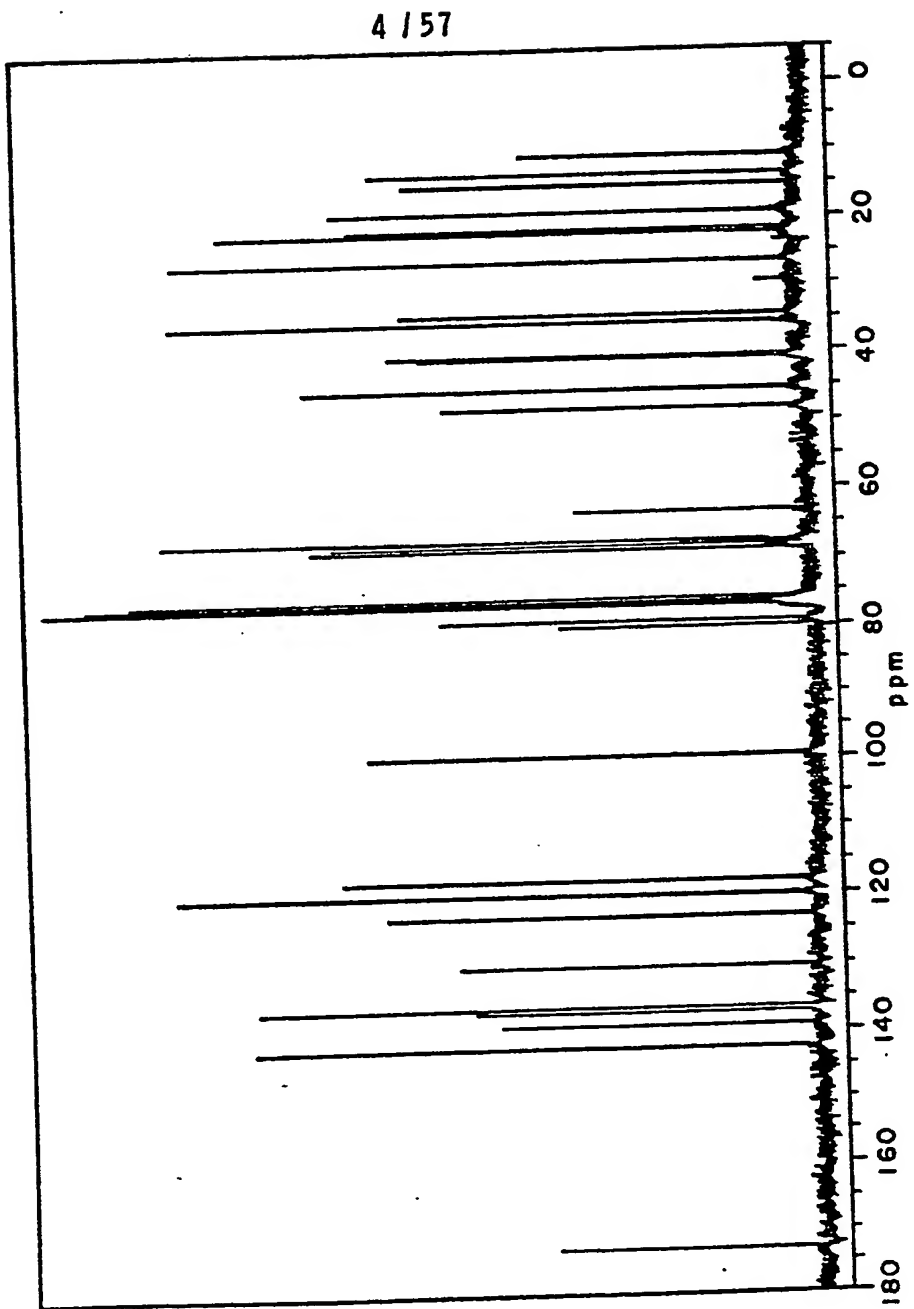
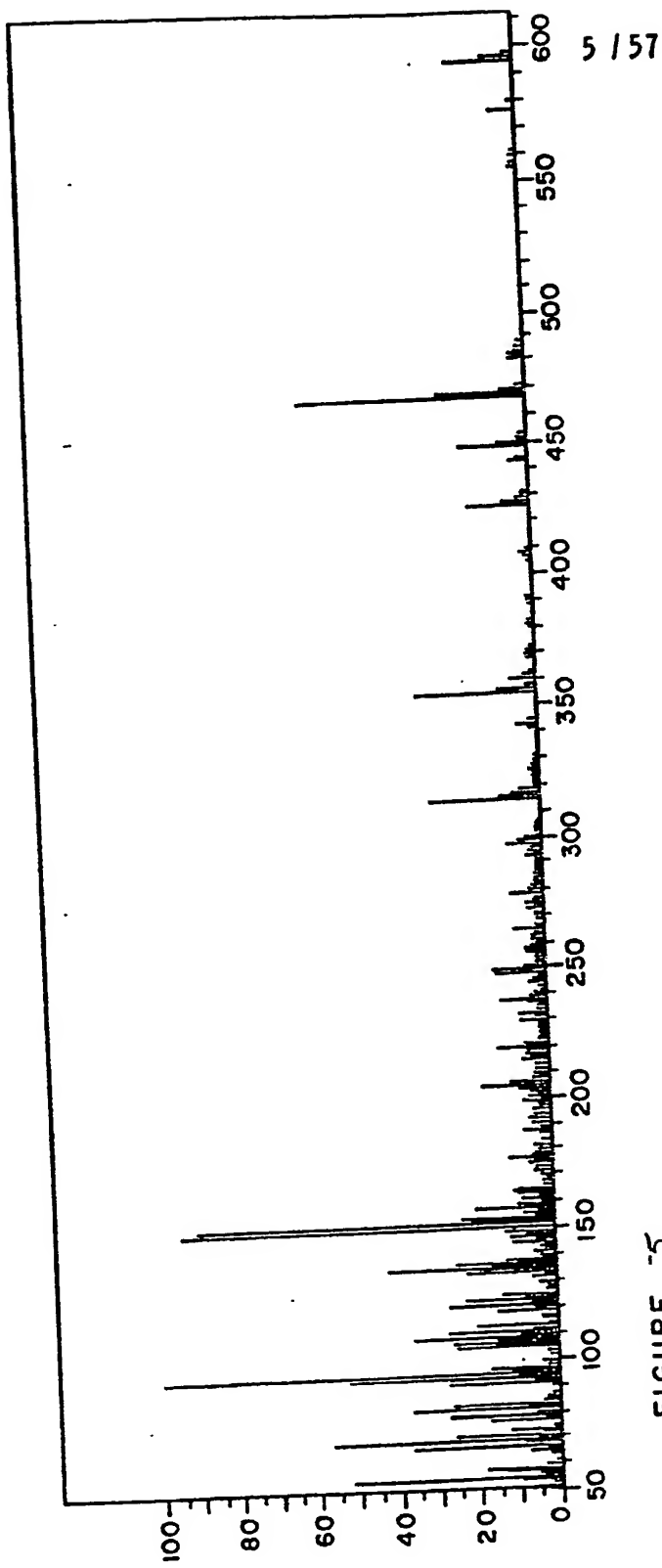


FIGURE 4

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ELECTRON IMPACT MASS SPECTRUM
OF LL-F28249α



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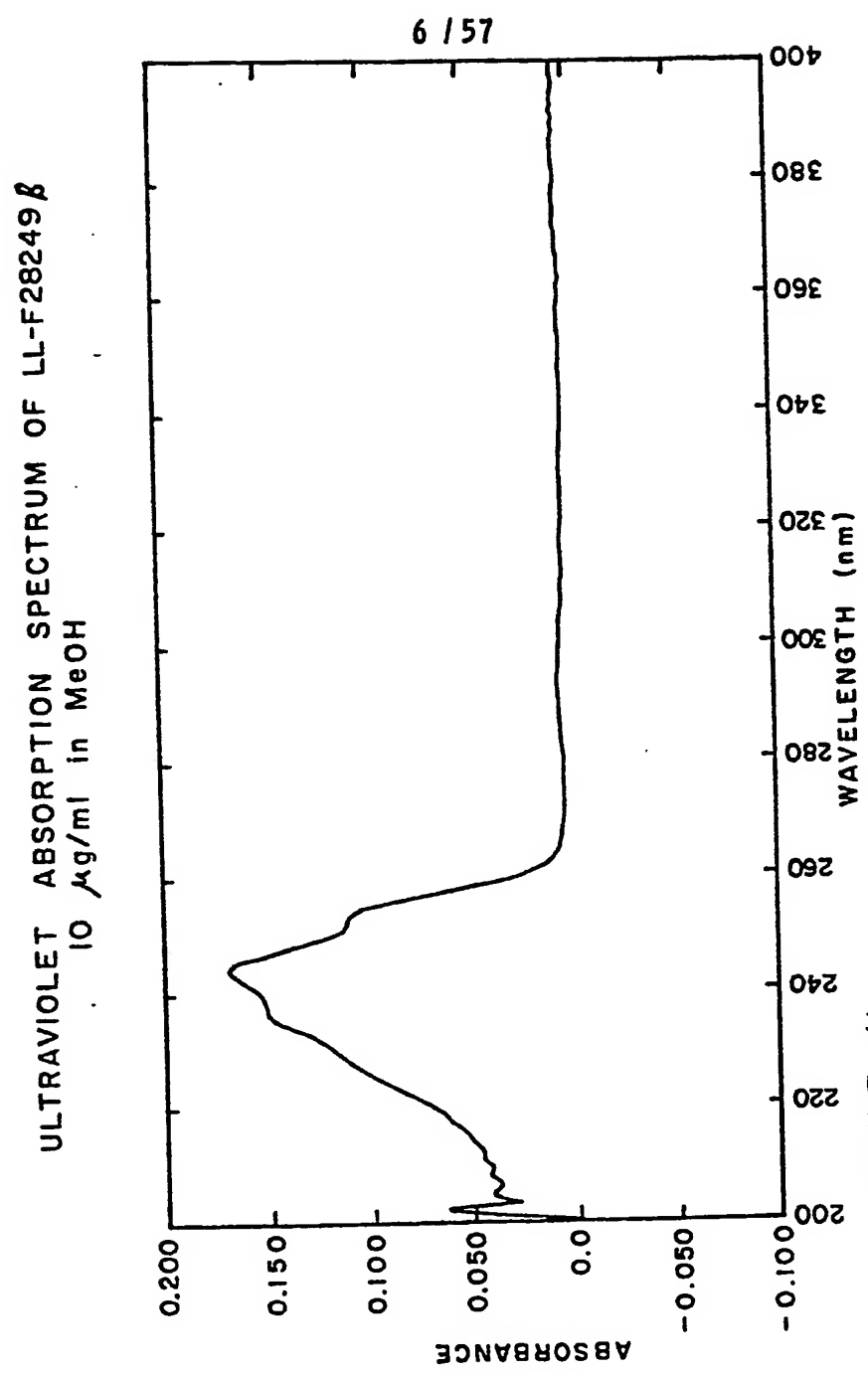


FIGURE 6.

0170006

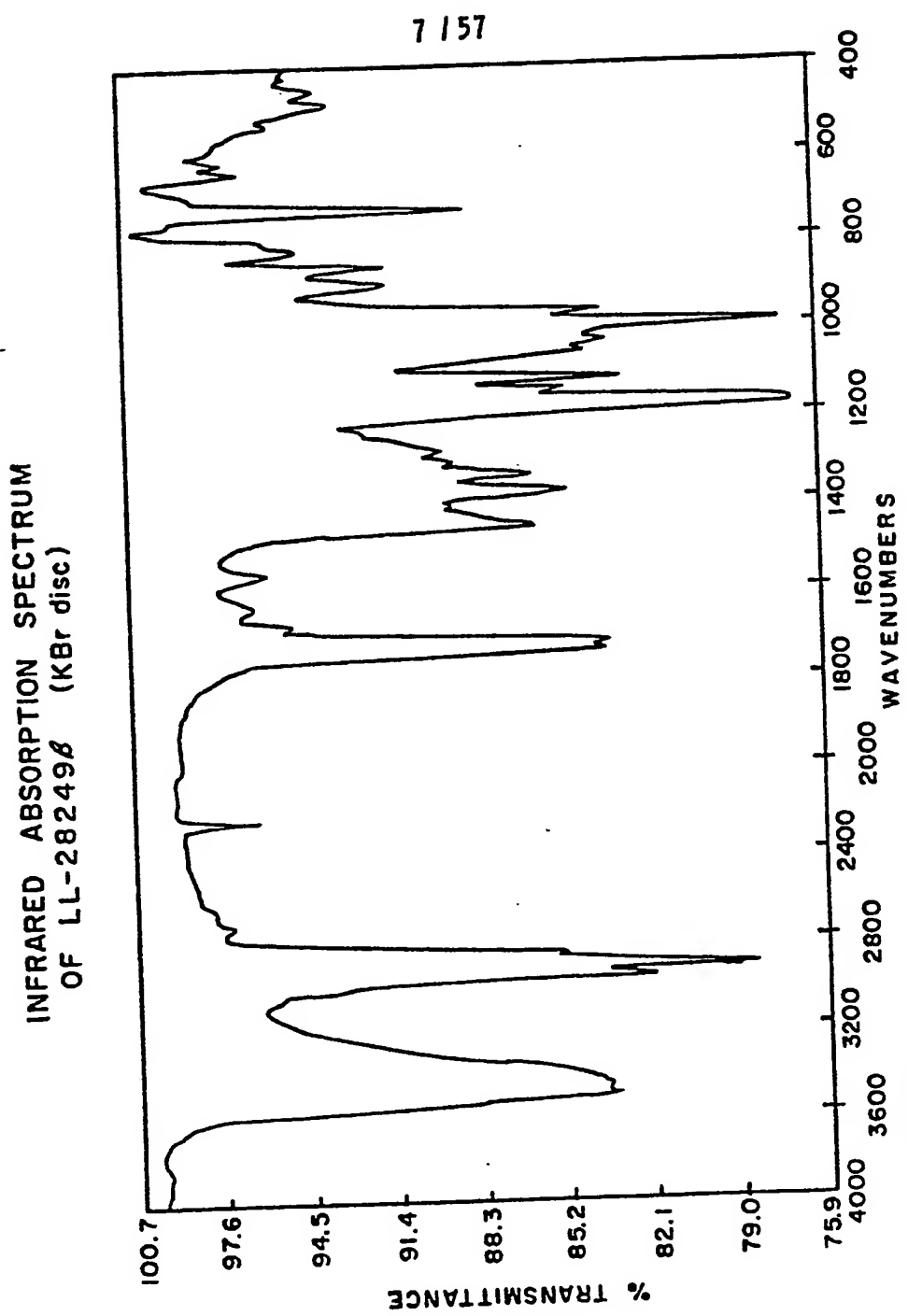


FIGURE 7

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PROTON NUCLEAR MAGNETIC RESONANCE SPECTRUM
OF LL-F28249 β IN CDCl₃ SOLUTION

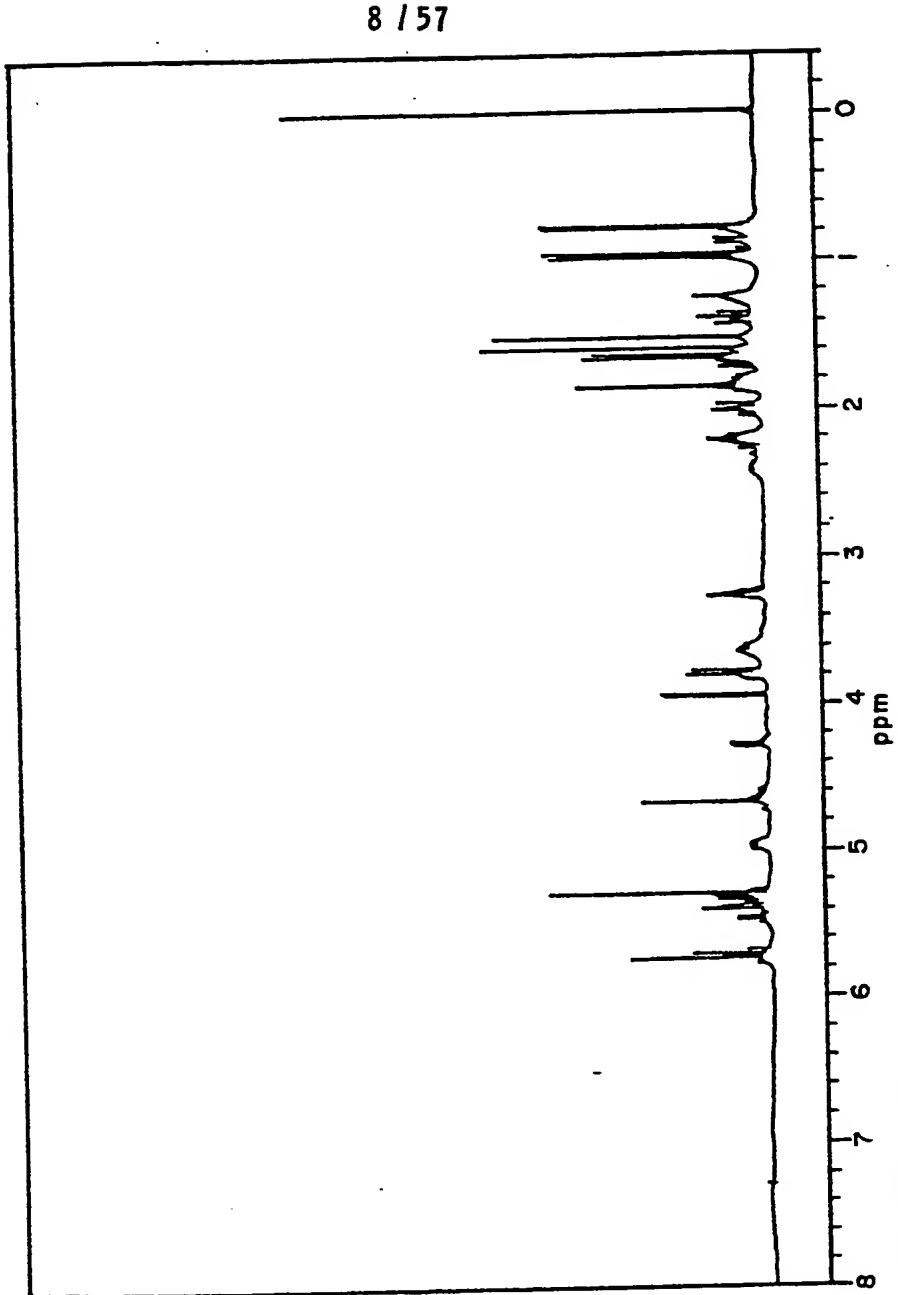


FIGURE 8

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ELECTRON IMPACT MASS SPECTRUM OF LL-F28249 β

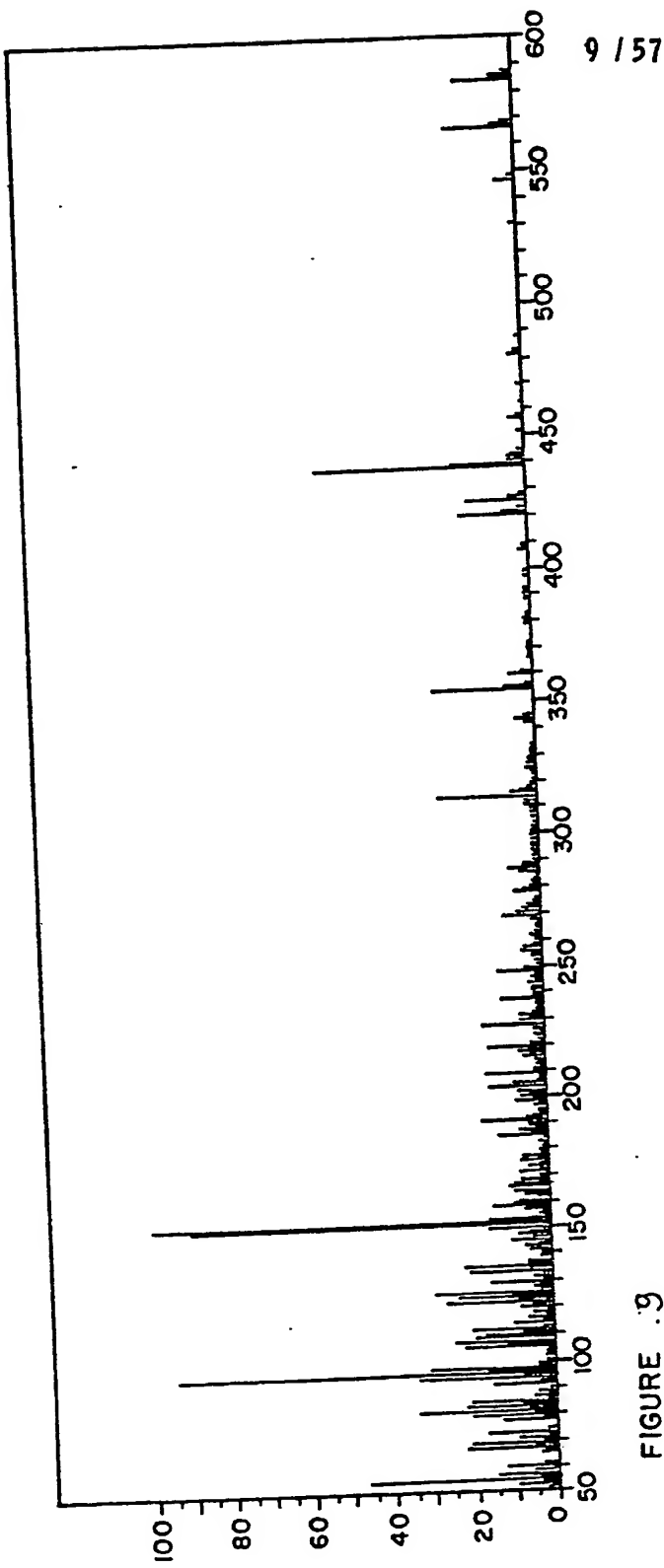


FIGURE .9

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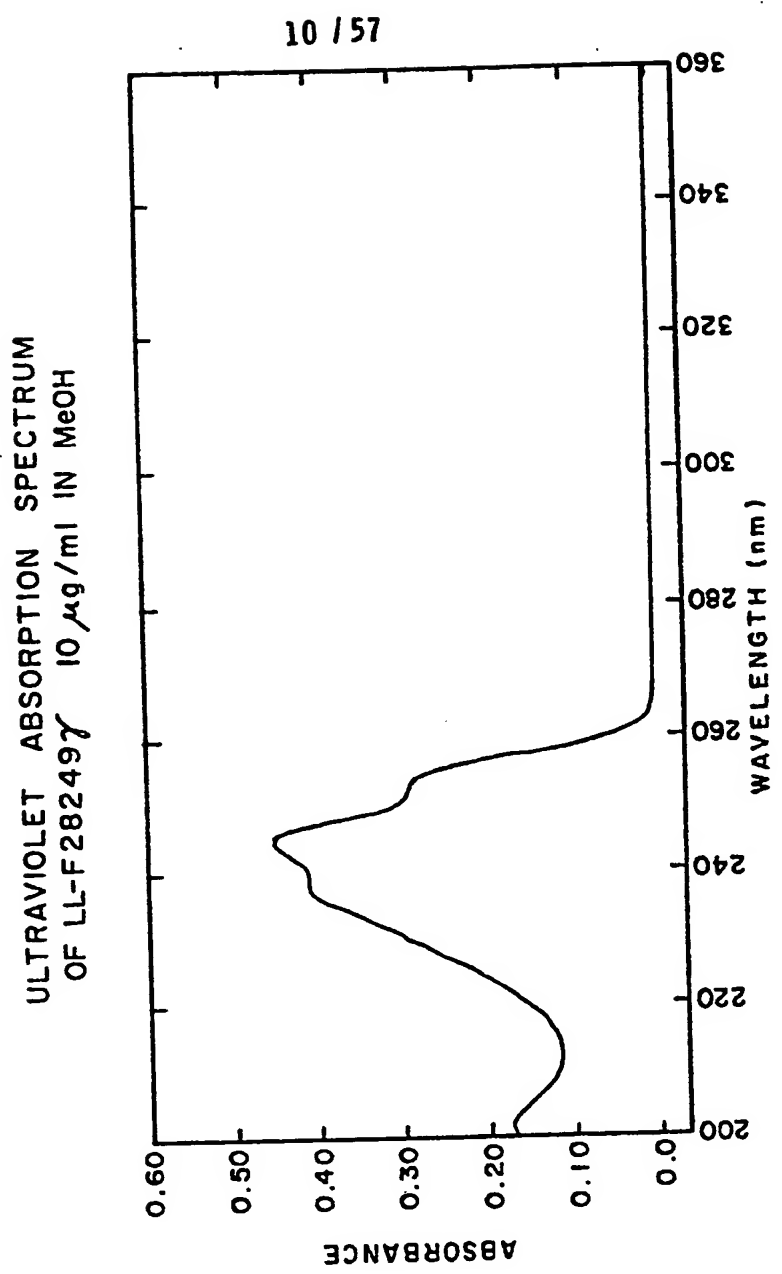


FIGURE 10

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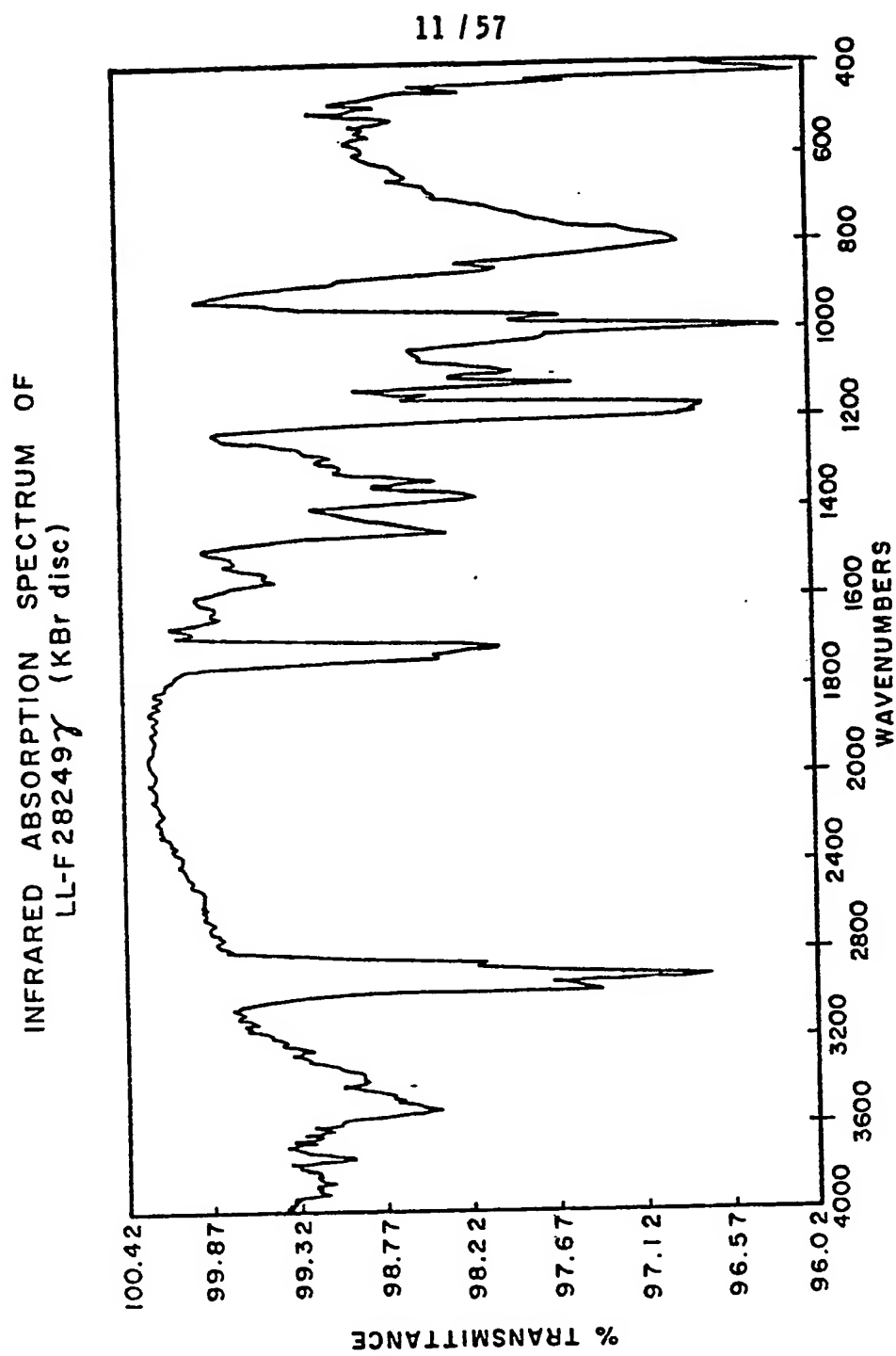


FIGURE 11

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PROTON NUCLEAR MAGNETIC RESONANCE SPECTRUM
OF LL-F28249γ IN CDCl₃ SOLUTION

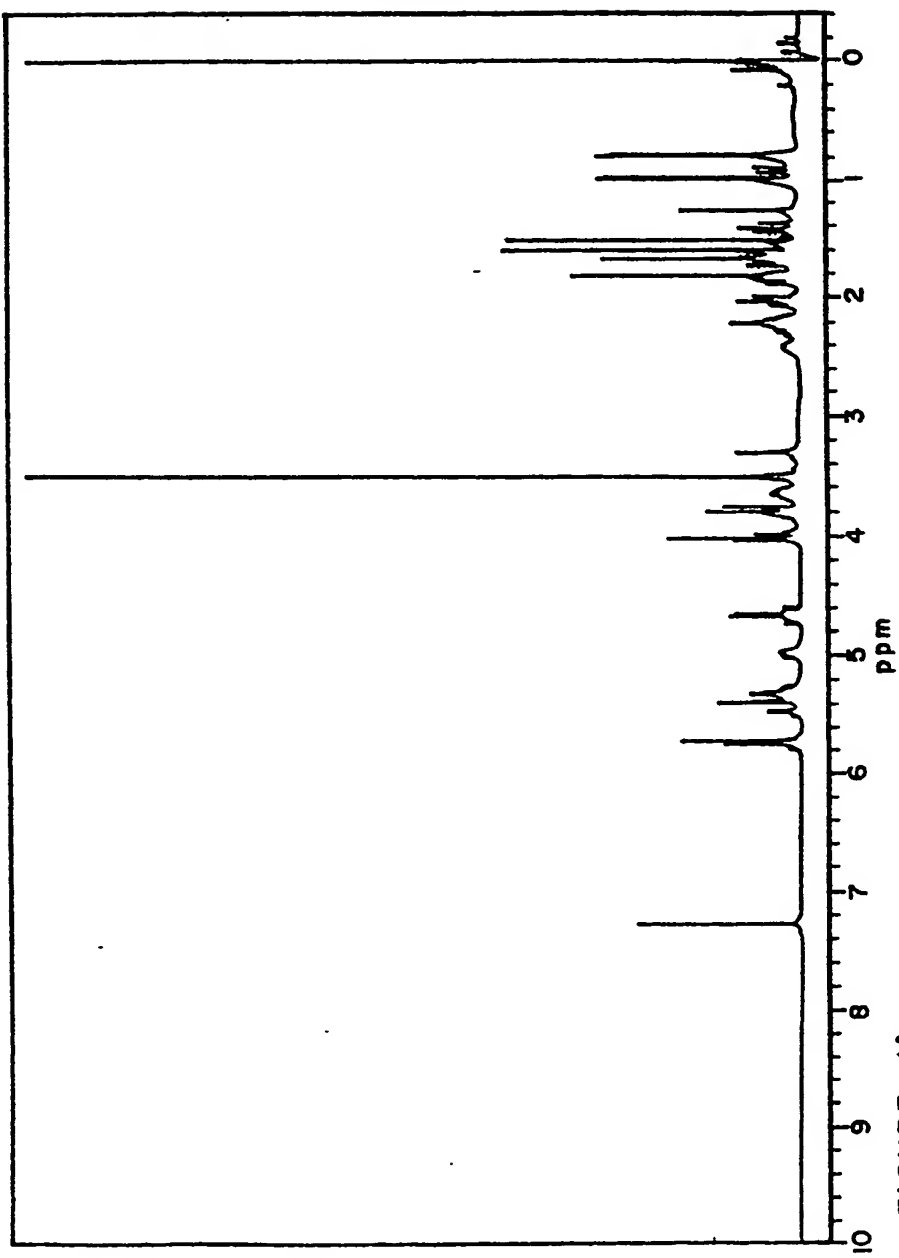


FIGURE 12

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CARBON-13 NUCLEAR MAGNETIC RESONANCE SPECTRUM
OF LL-F28249γ IN CDCl₃ SOLUTION

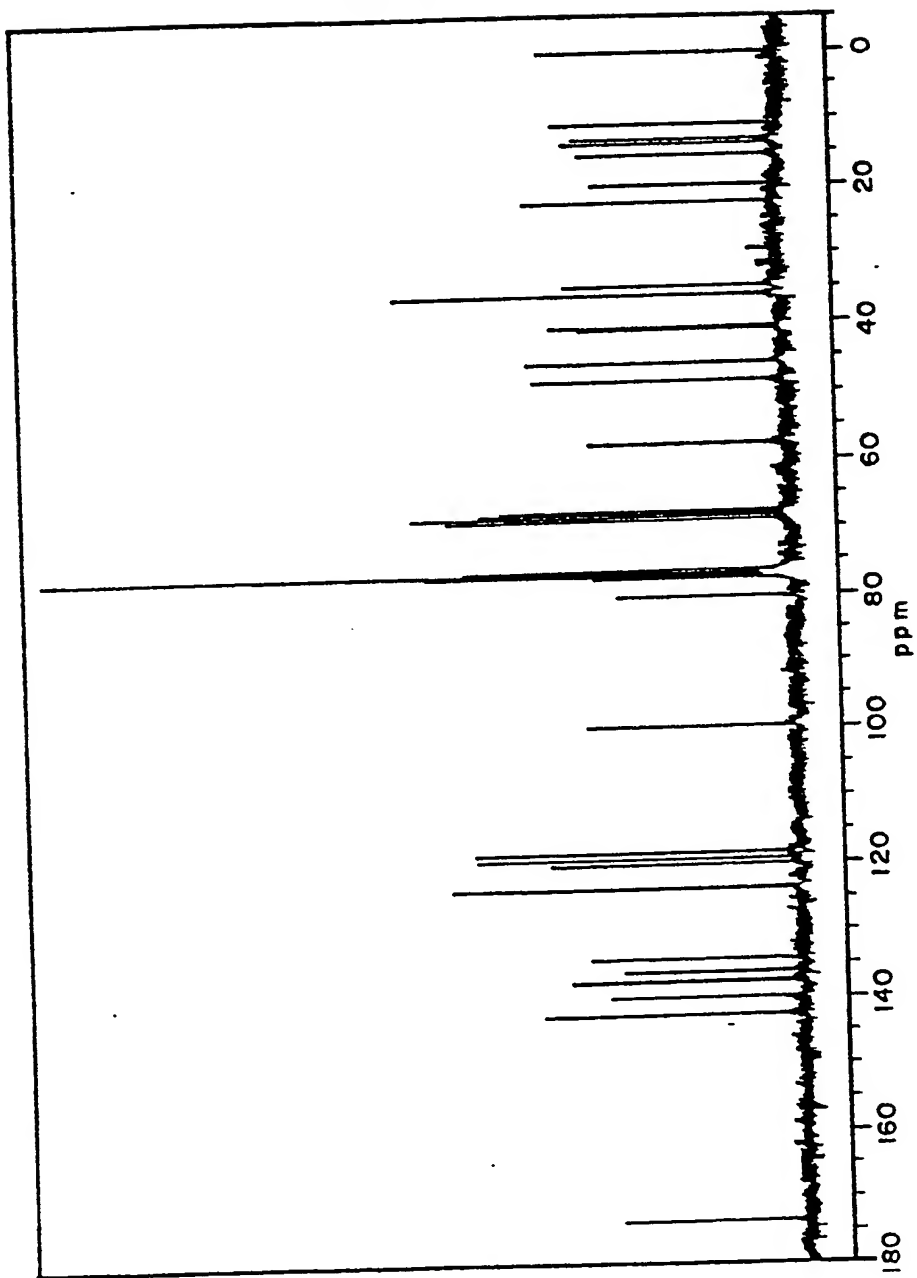


FIGURE 13

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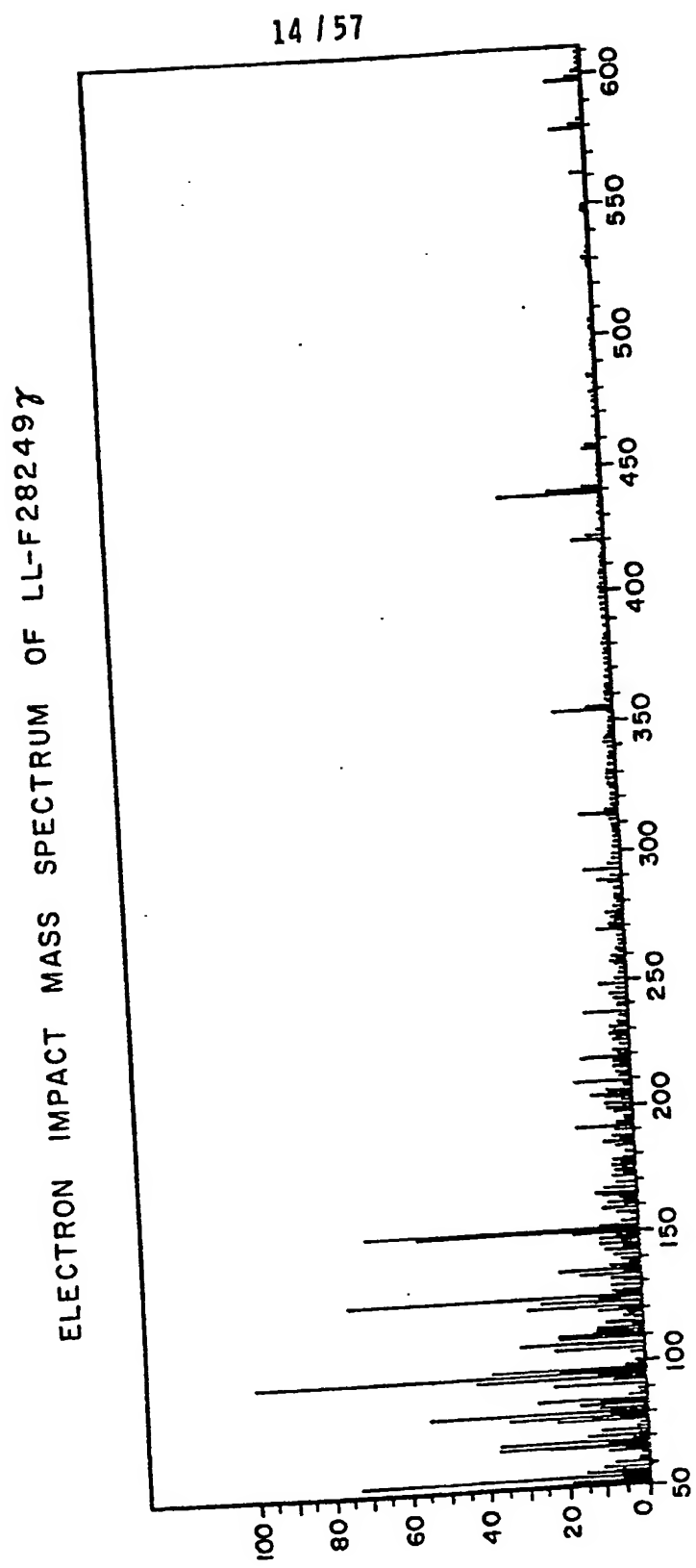


FIGURE A4

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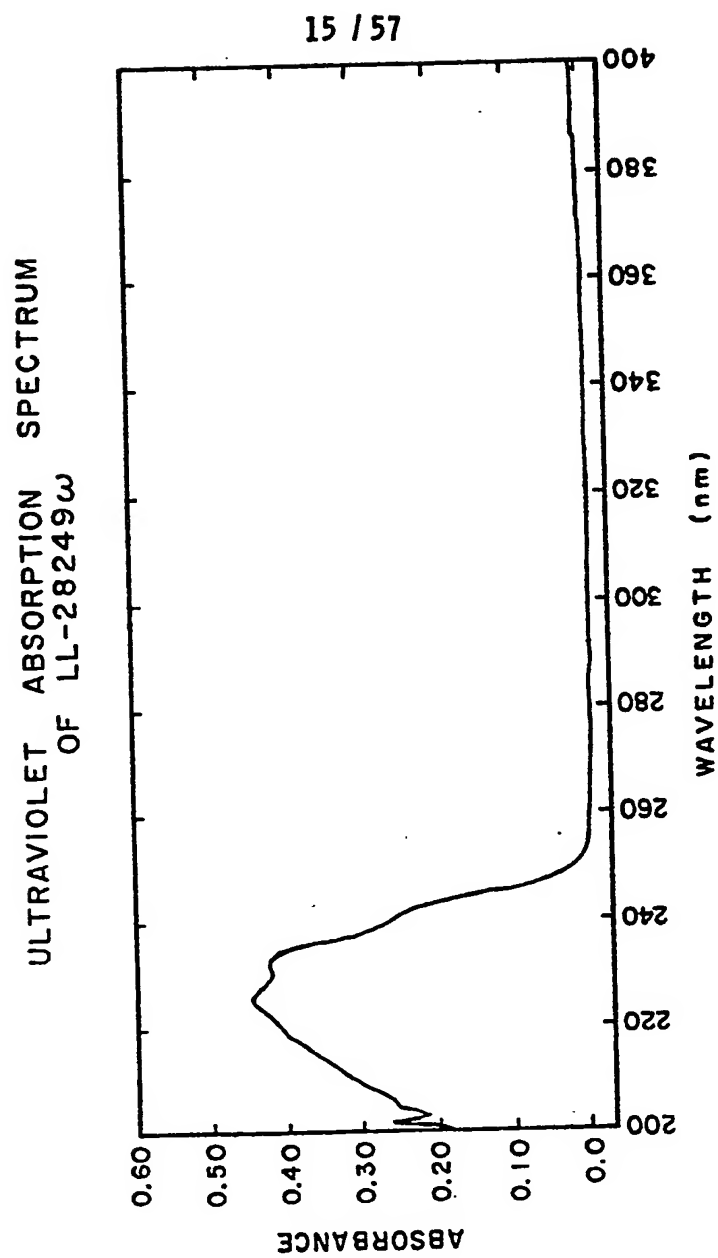


FIGURE 45

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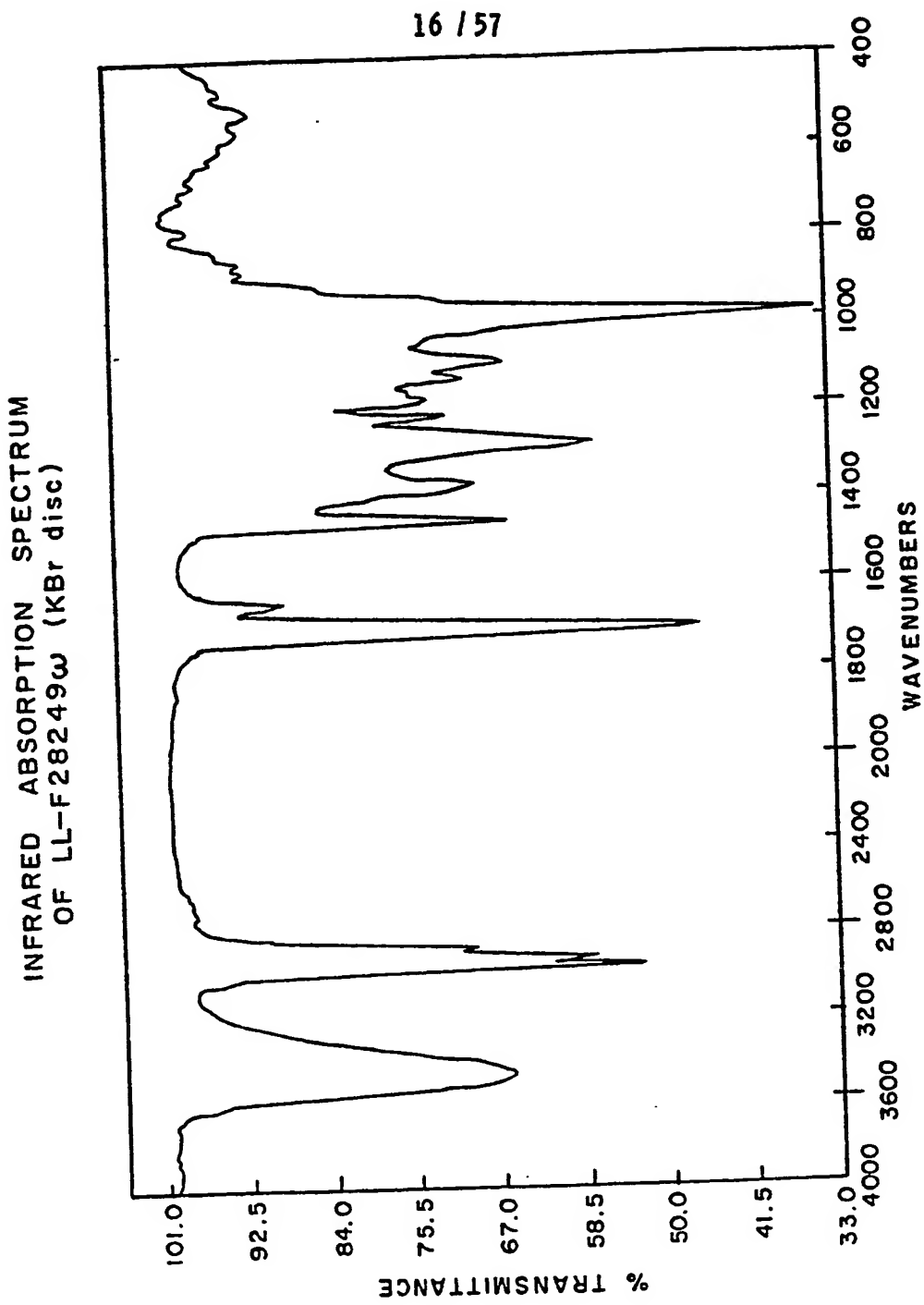


FIGURE 16

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PROTON NUCLEAR MAGNETIC RESONANCE SPECTRUM
OF LL-F28249 ω IN CDCl₃ SOLUTION

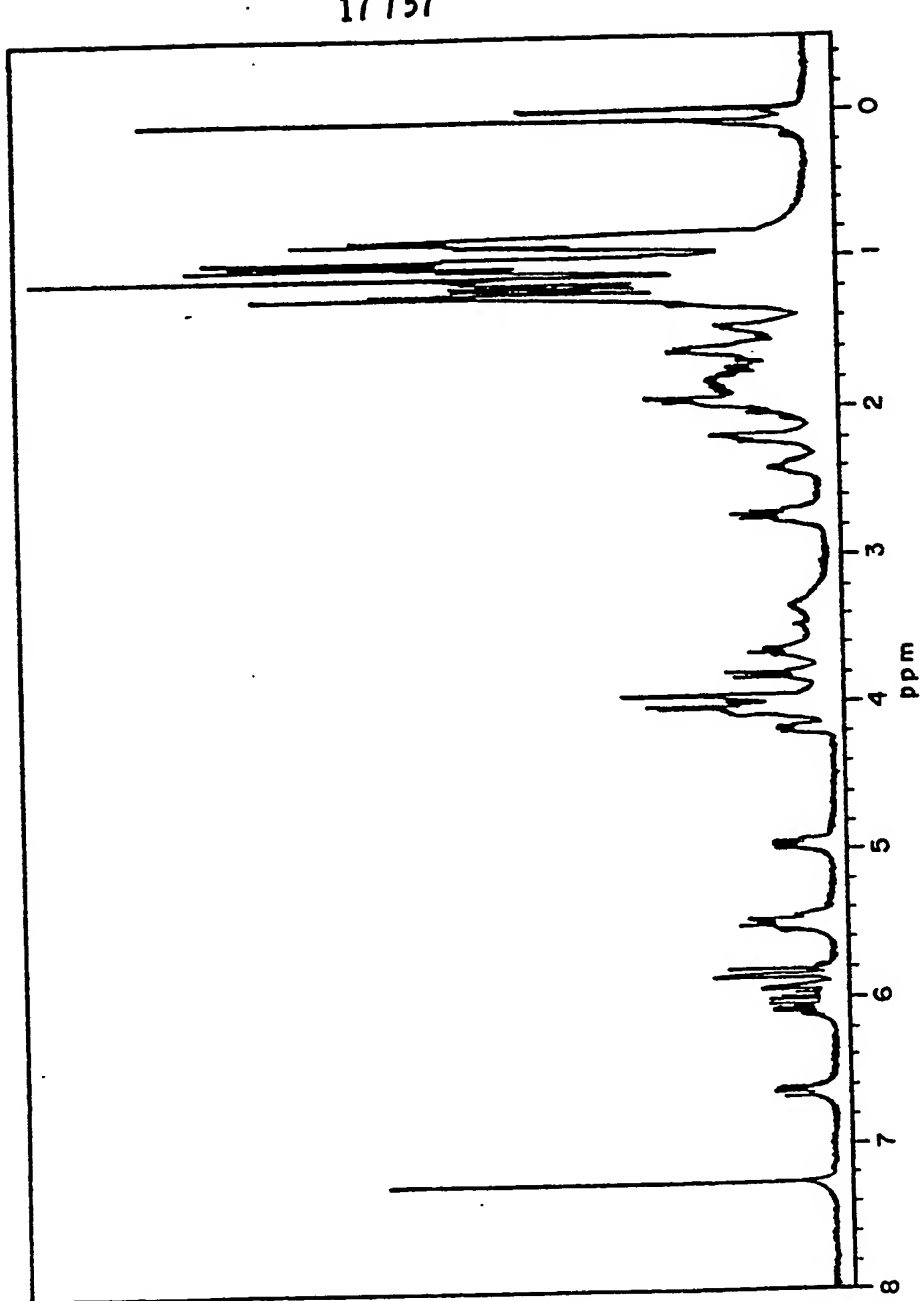


FIGURE . 17

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CARBON-13 NUCLEAR MAGNETIC RESONANCE SPECTRUM
OF LL-F28249W IN CDCl₃ SOLUTION

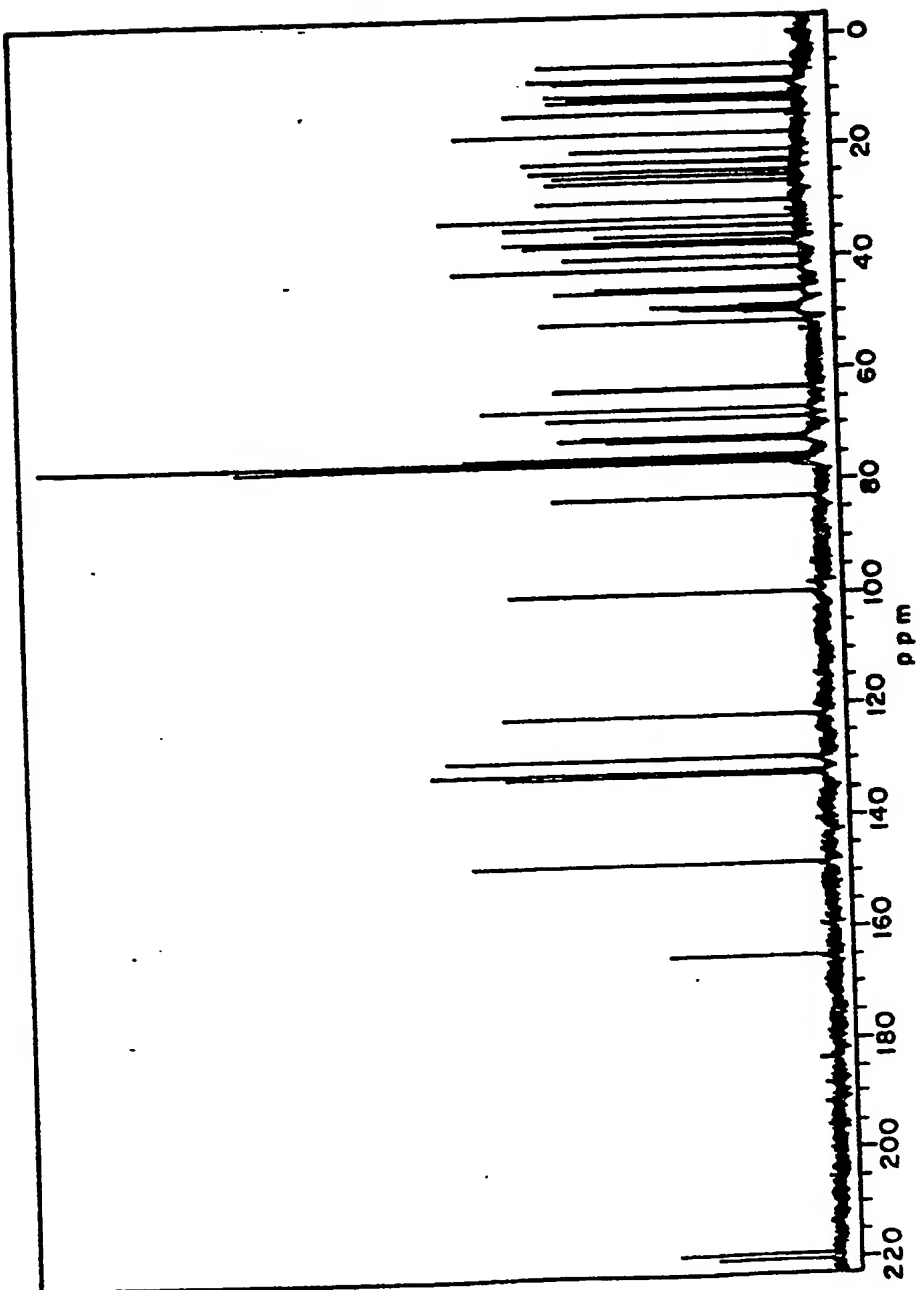


FIGURE 18.

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ELECTRON IMPACT MASS SPECTRUM
OF LL-F28249_u

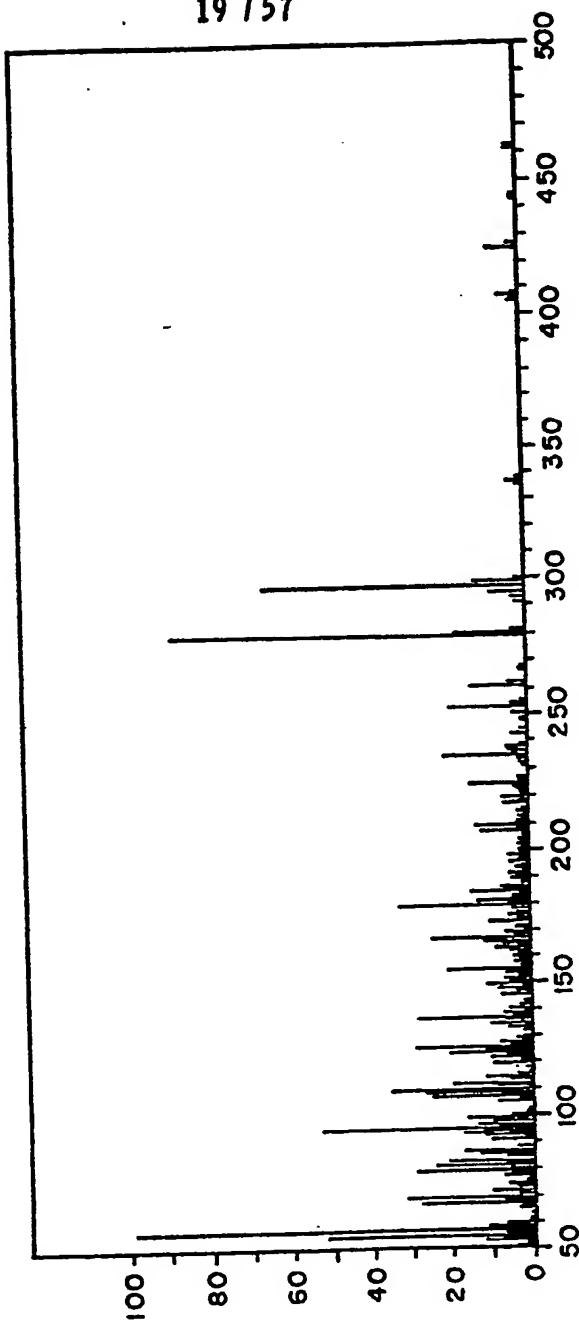
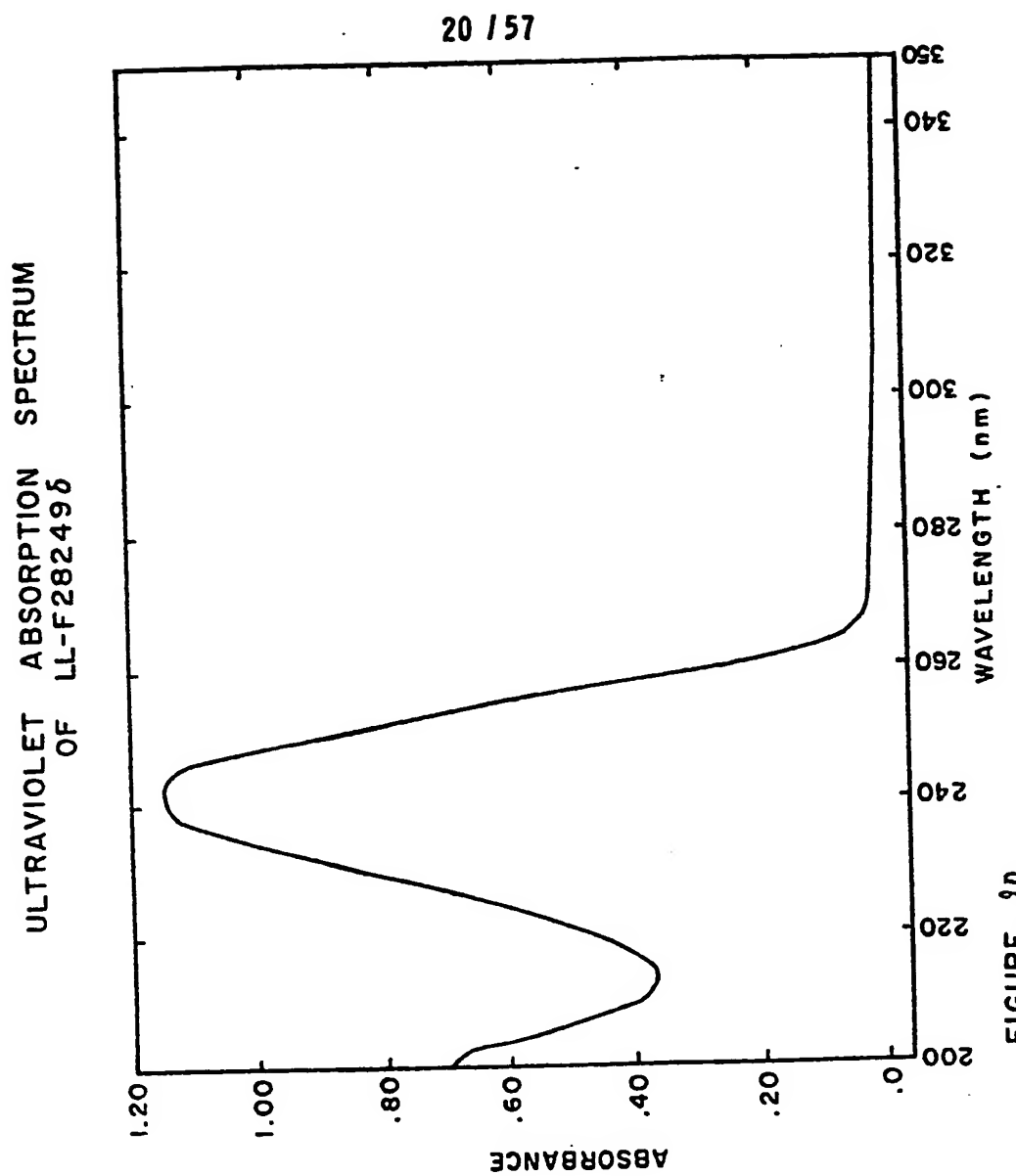


FIGURE -19

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PROTON NUCLEAR MAGNETIC RESONANCE SPECTRUM
OF LL-F28249 δ IN CDCl₃ SOLUTION

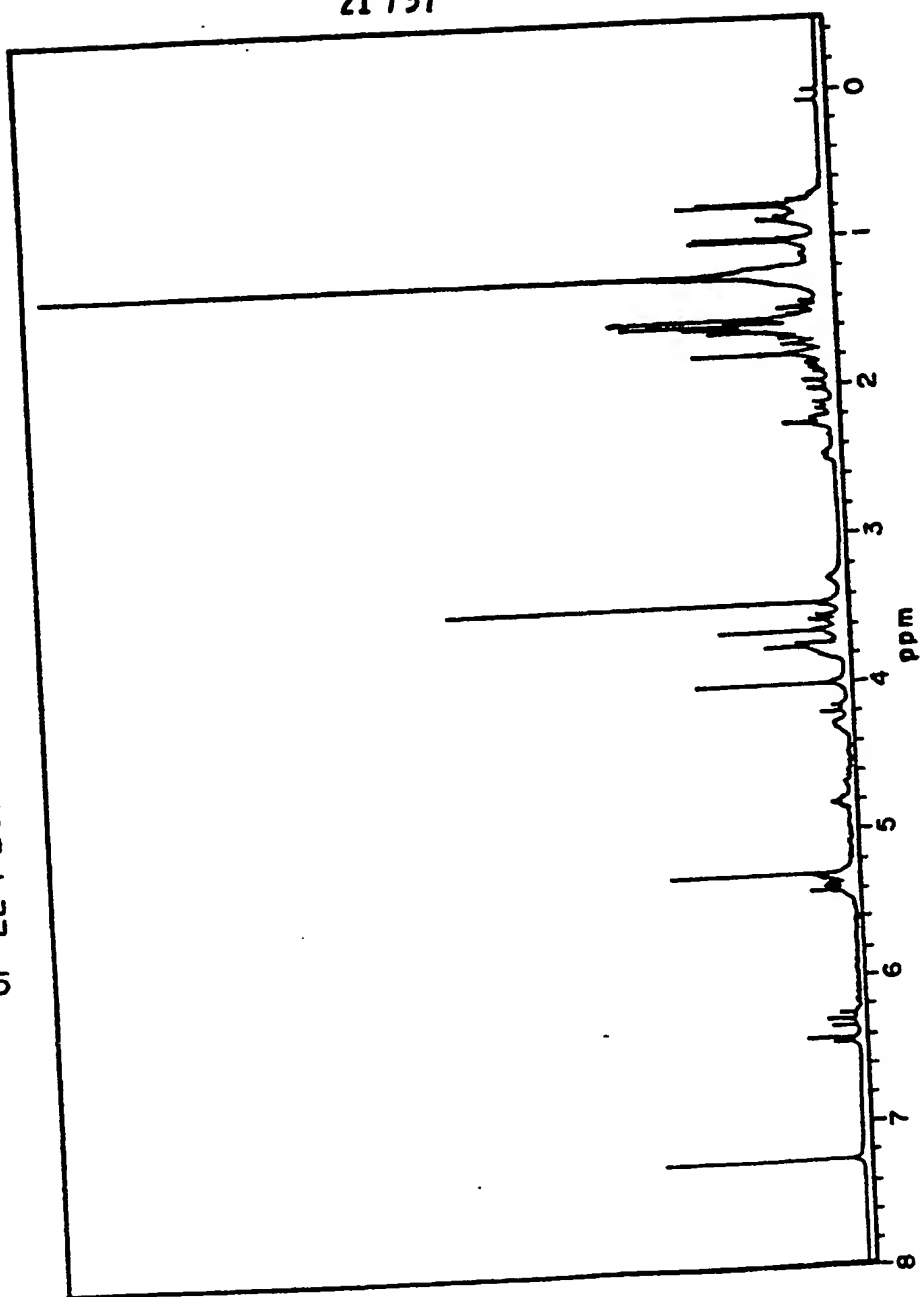


FIGURE 21

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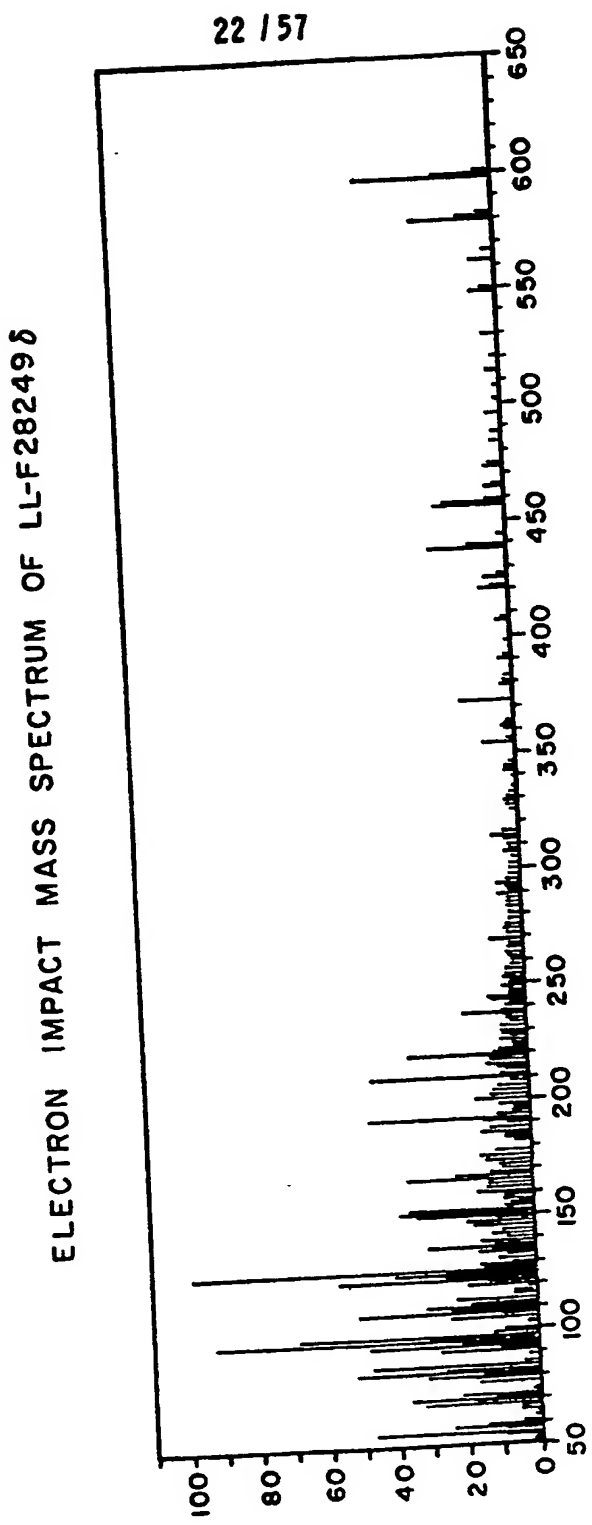
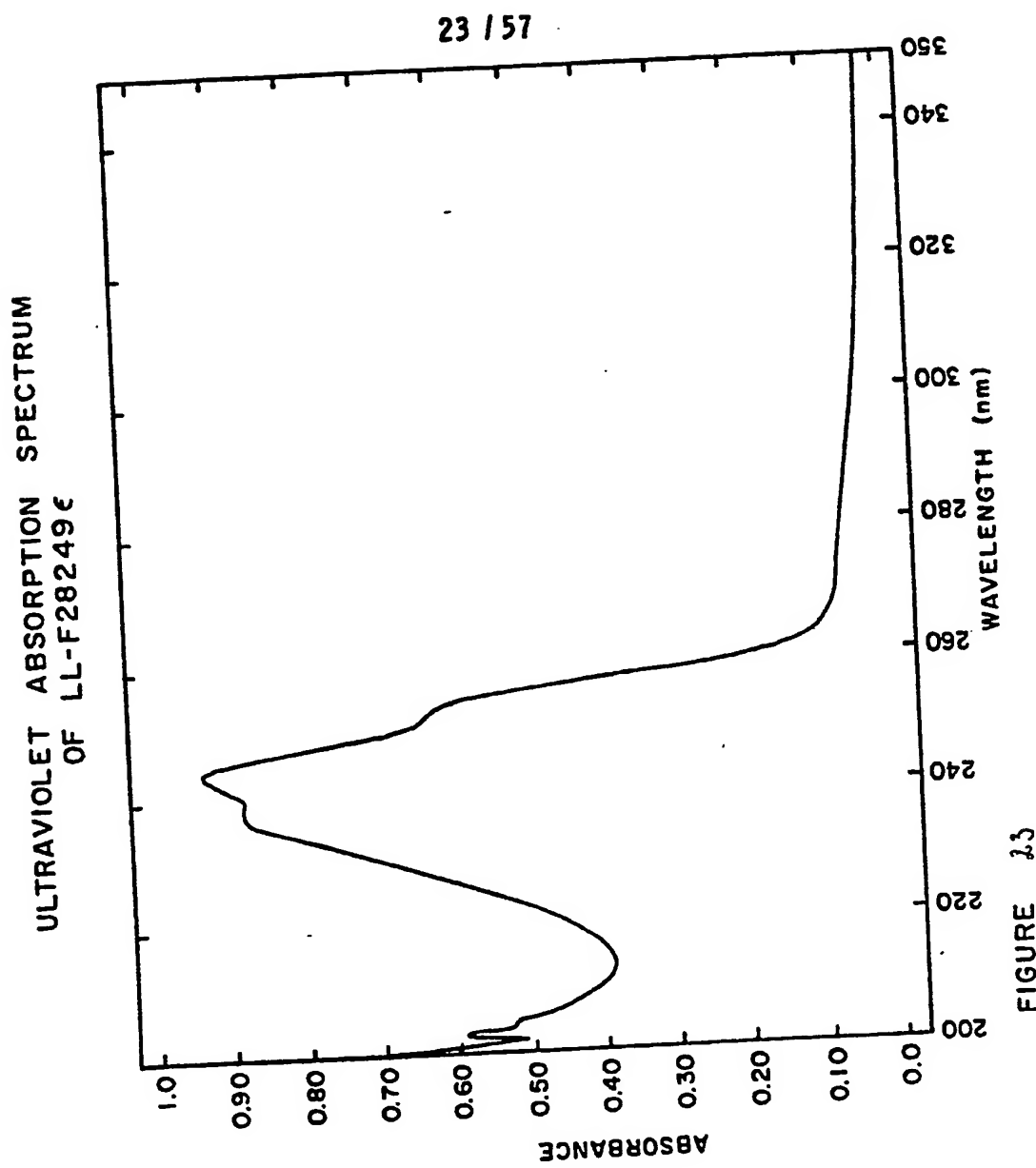


FIGURE 2:

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PROTON NUCLEAR MAGNETIC RESONANCE SPECTRUM
OF LL-F28249c IN CDCl₃ SOLUTION

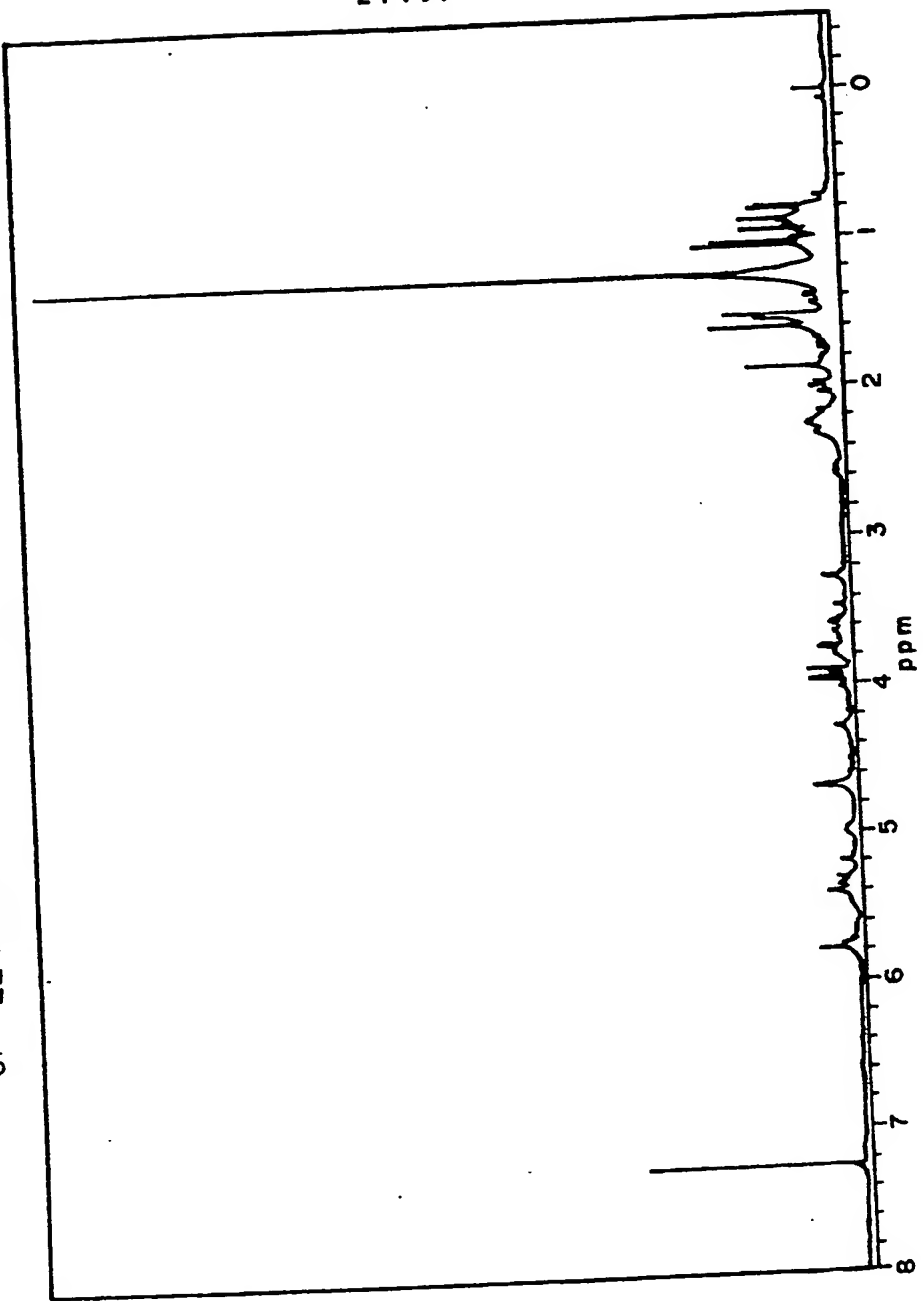


FIGURE 24

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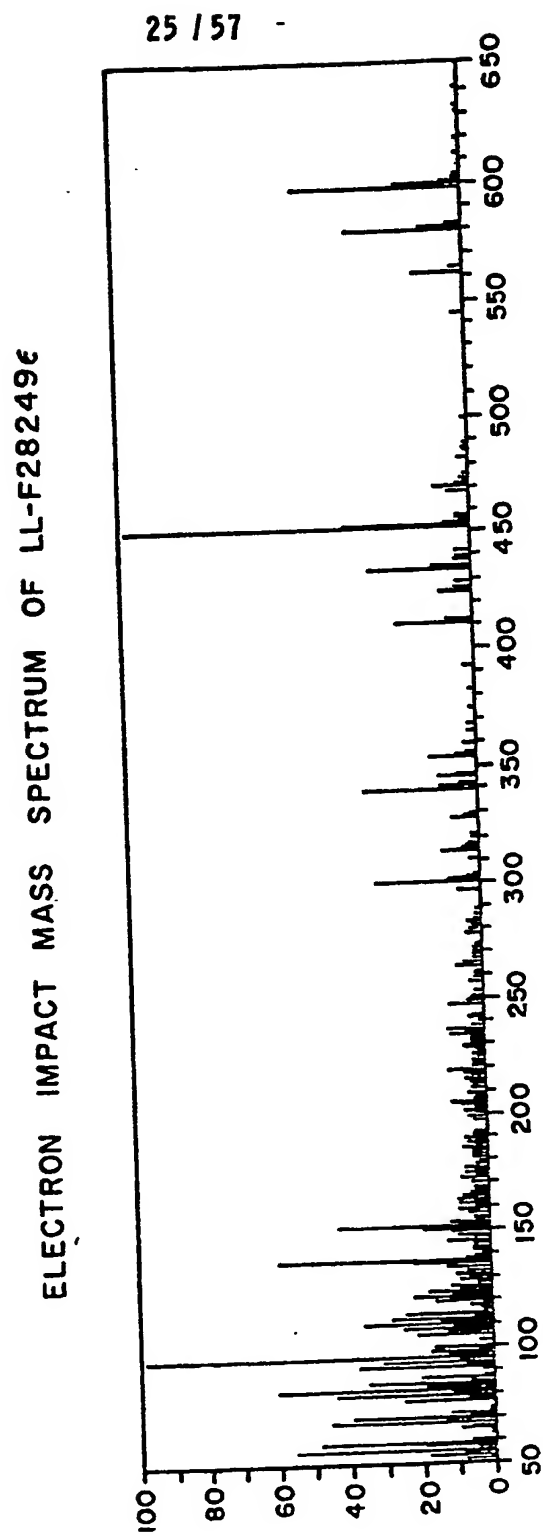


FIGURE 25

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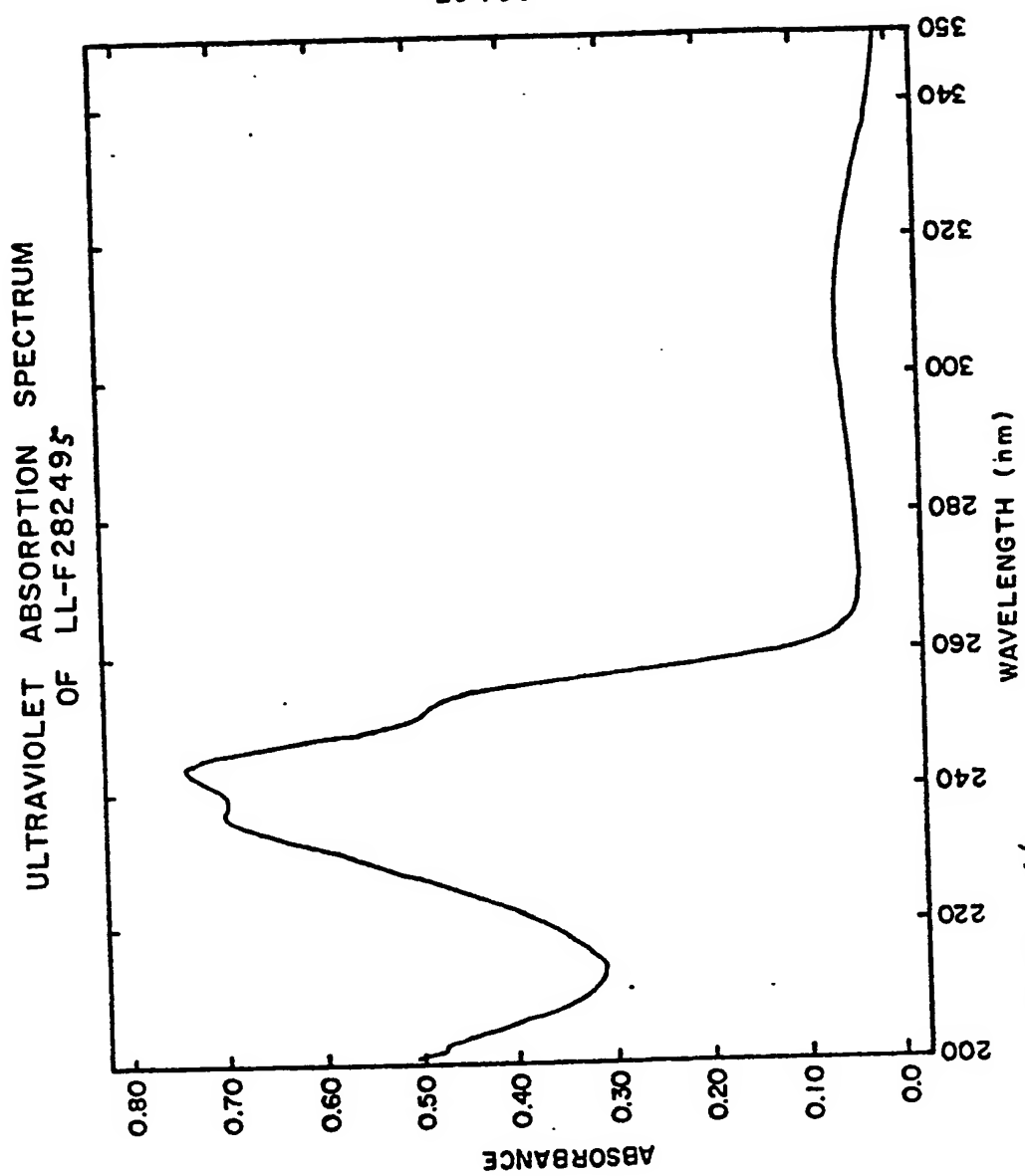


FIGURE . 26

0170006

PROTON NUCLEAR MAGNETIC RESONANCE SPECTRUM OF
LL-F282495 IN CDCl_3 SOLUTION

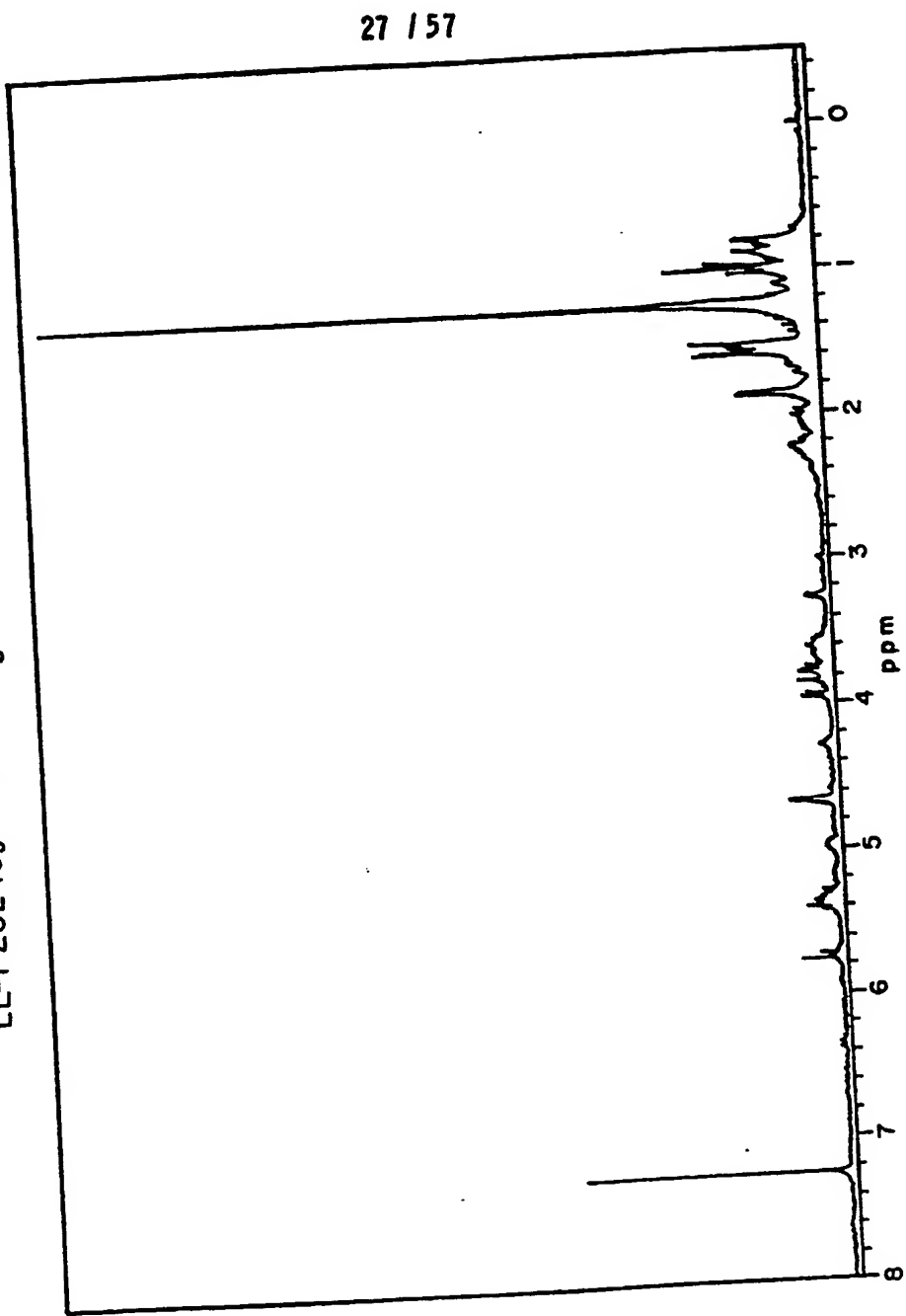


FIGURE 27

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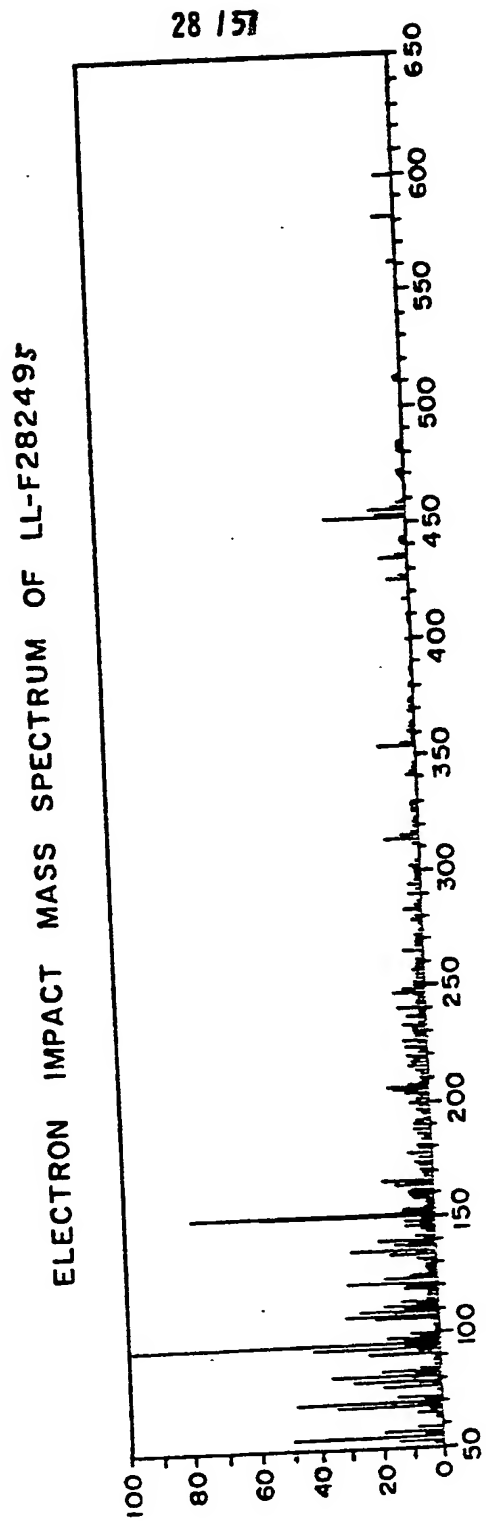


FIGURE 28

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ULTRAVIOLET ABSORPTION SPECTRUM OF LL-F282497

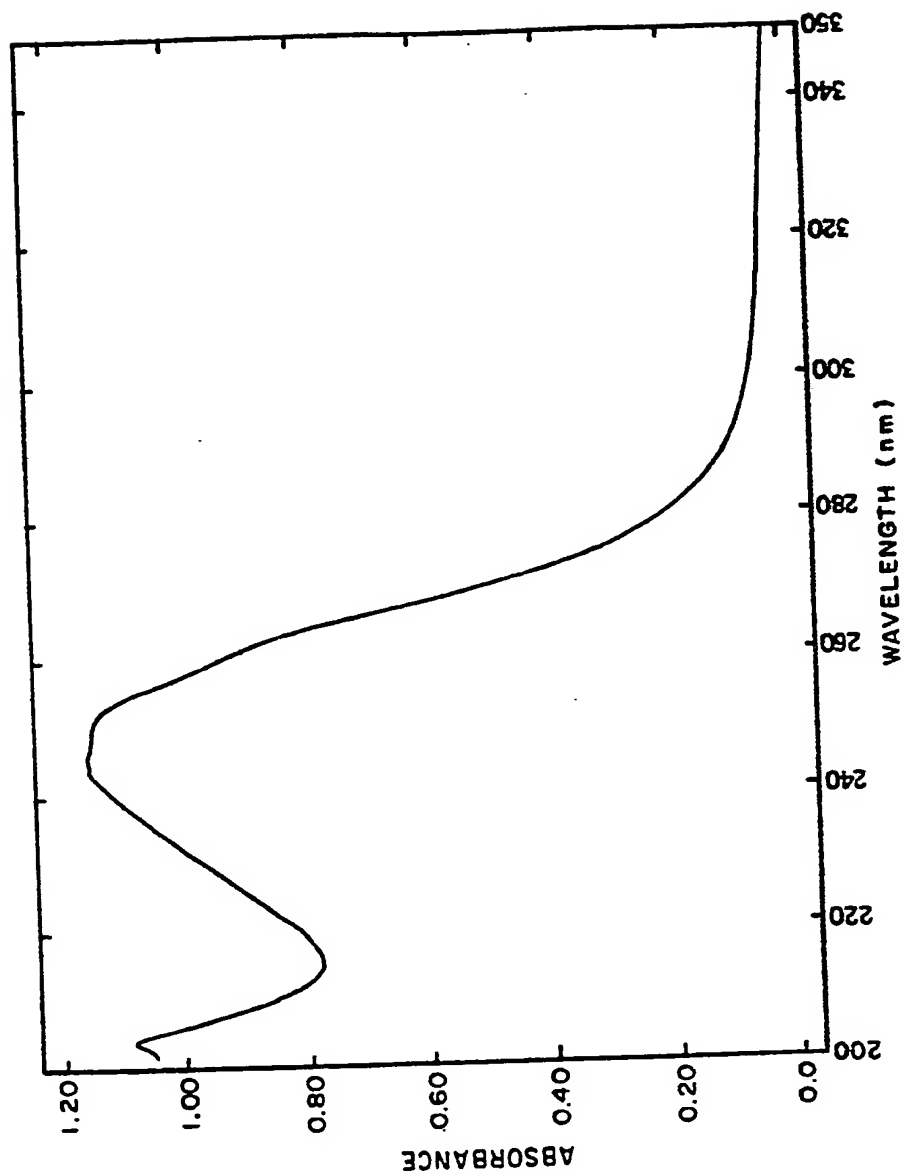


FIGURE . 19

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PROTON NUCLEAR MAGNETIC RESONANCE SPECTRUM OF
LL-F28249 η IN CDCl₃ SOLUTION

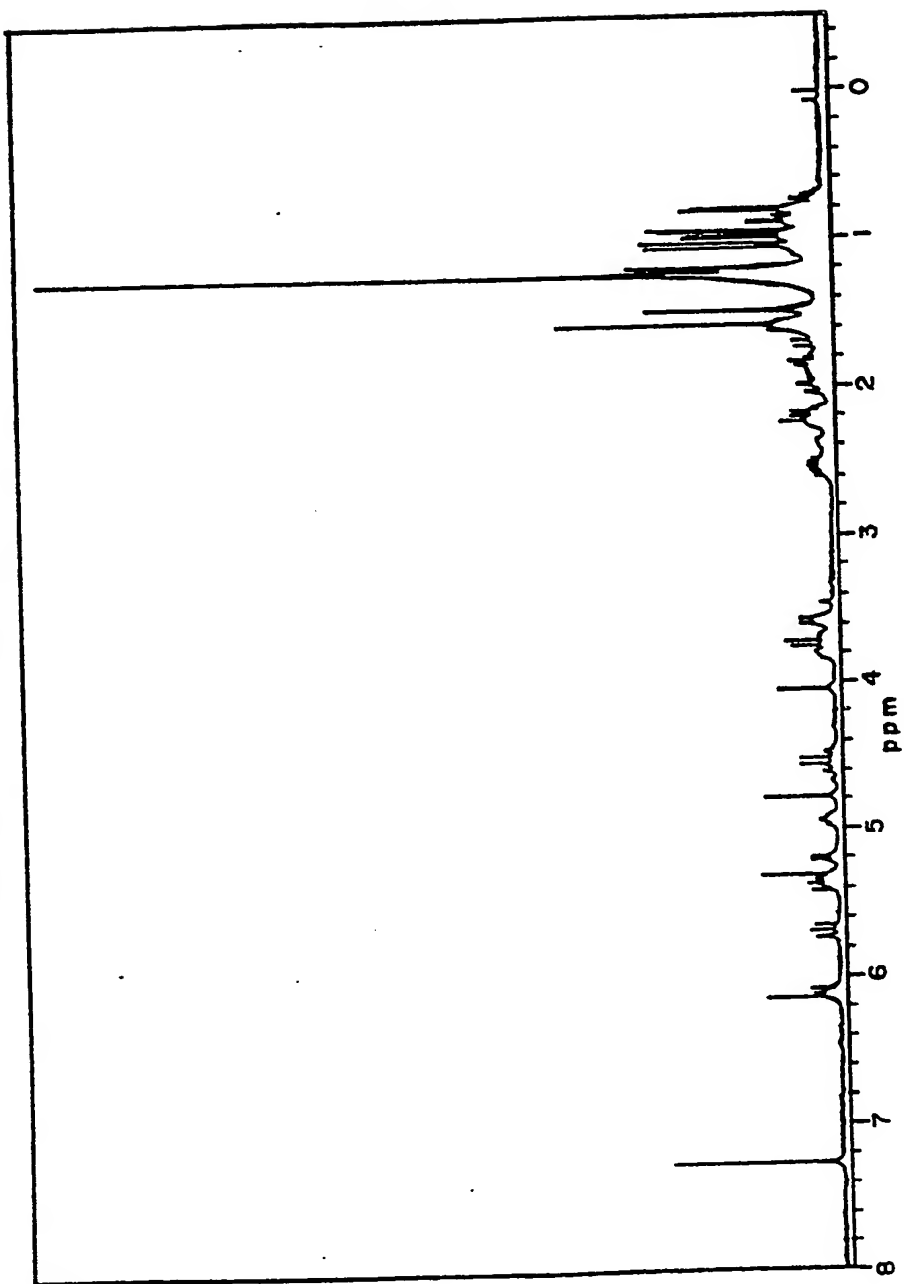


FIGURE 30.

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ELECTRON IMPACT MASS SPECTRUM OF LL-F282497

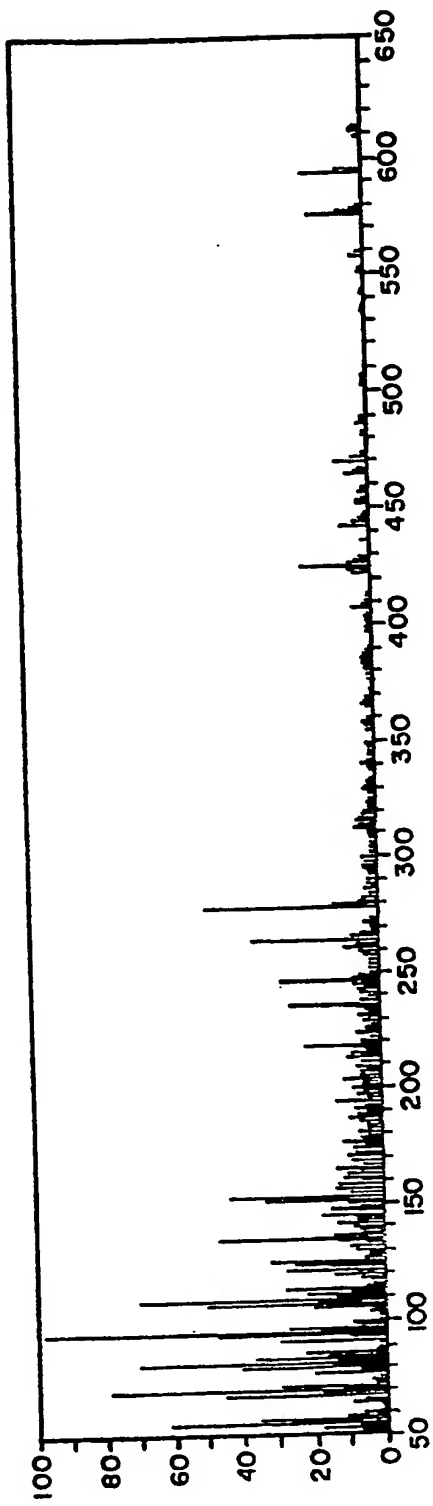


FIGURE . 31

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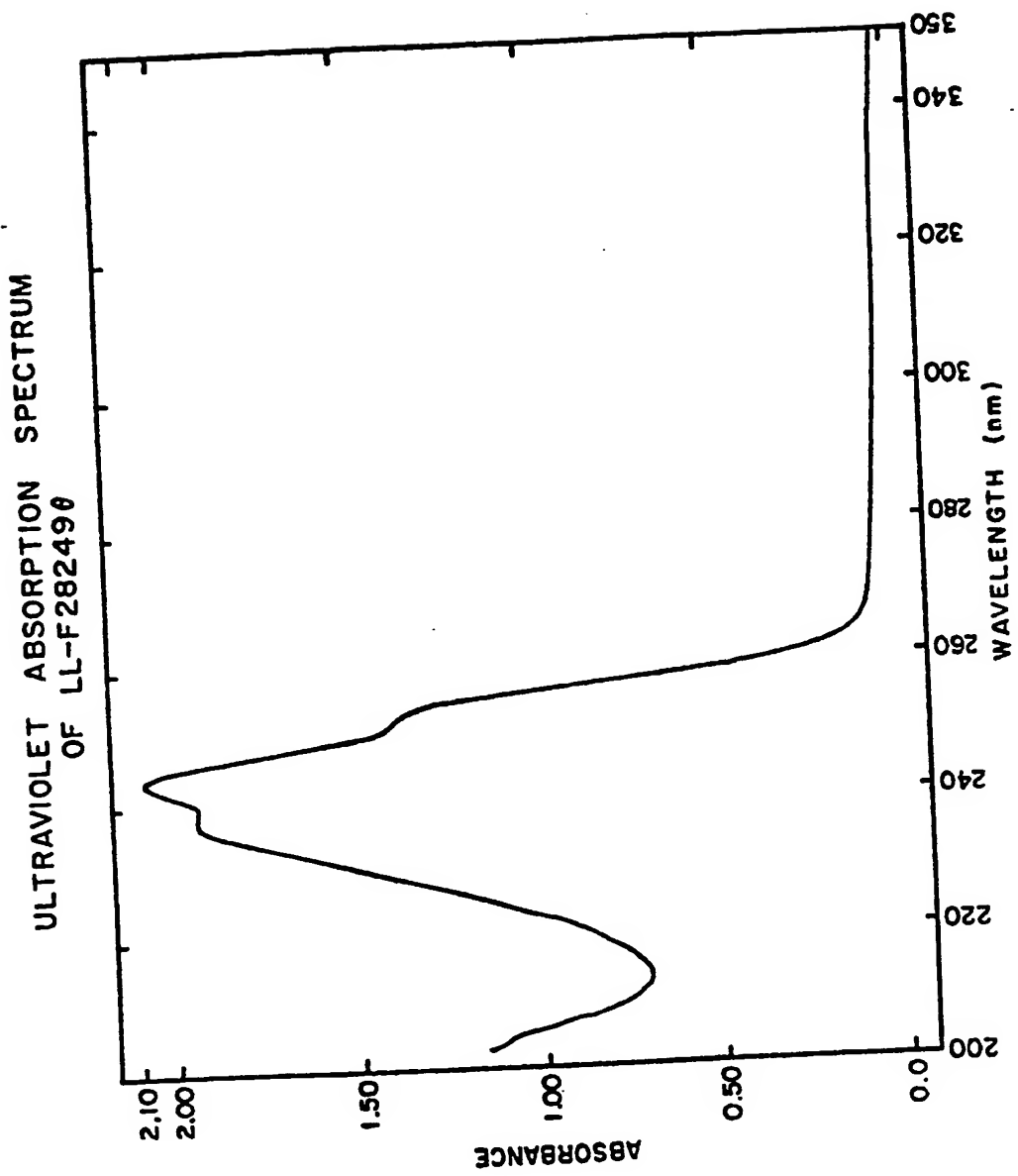


FIGURE 4

0170006

PROTON NUCLEAR MAGNETIC RESONANCE SPECTRUM OF
LL-F282490 IN CDCl₃ SOLUTION

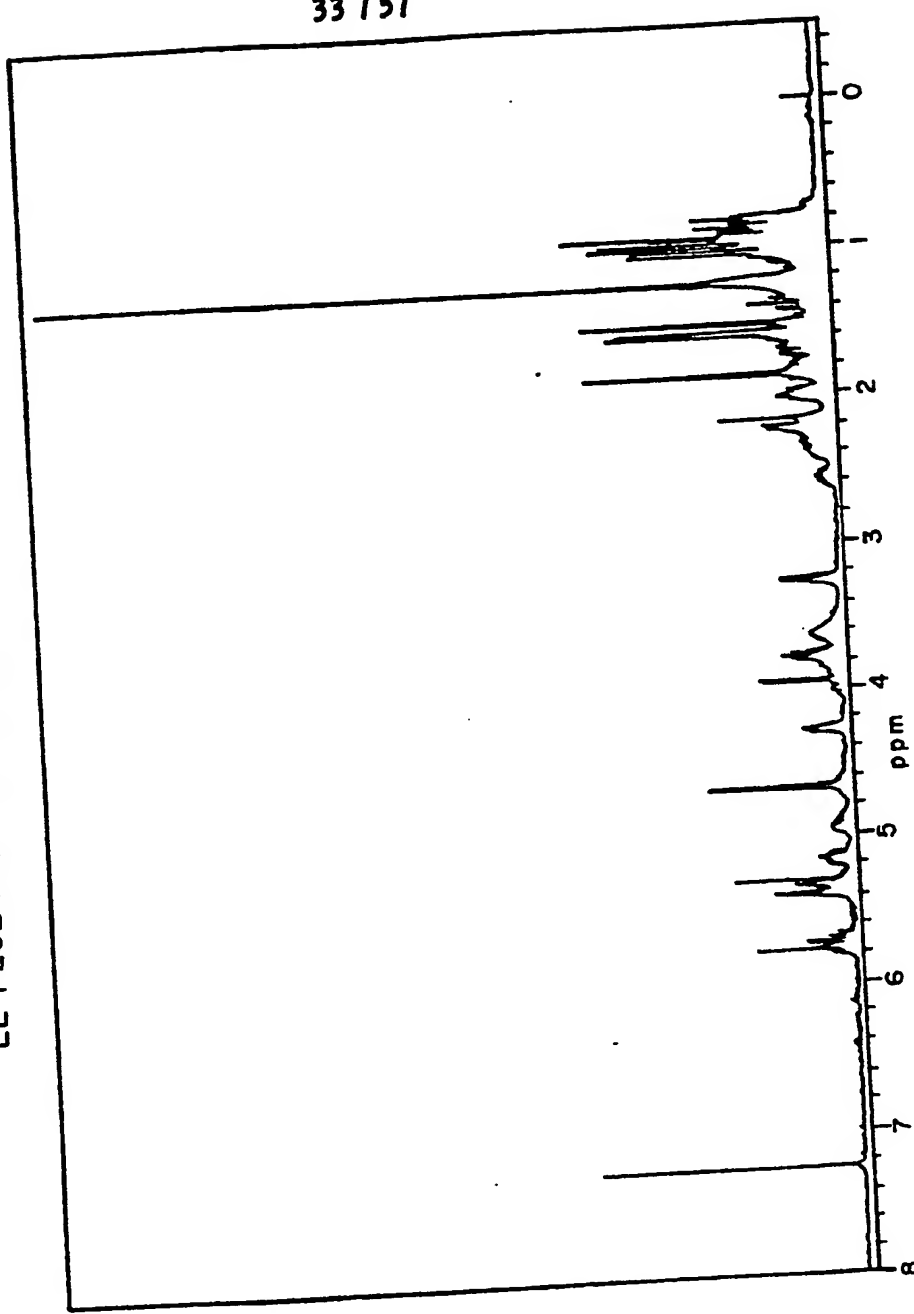


FIGURE 3j

0170006

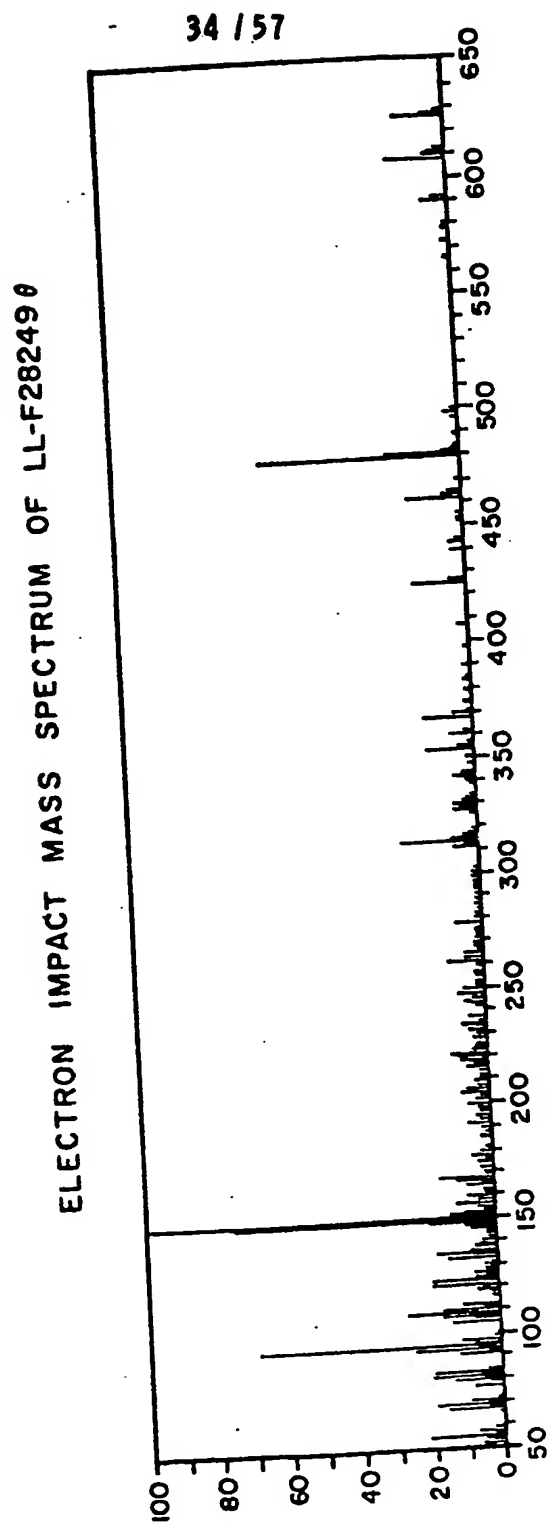
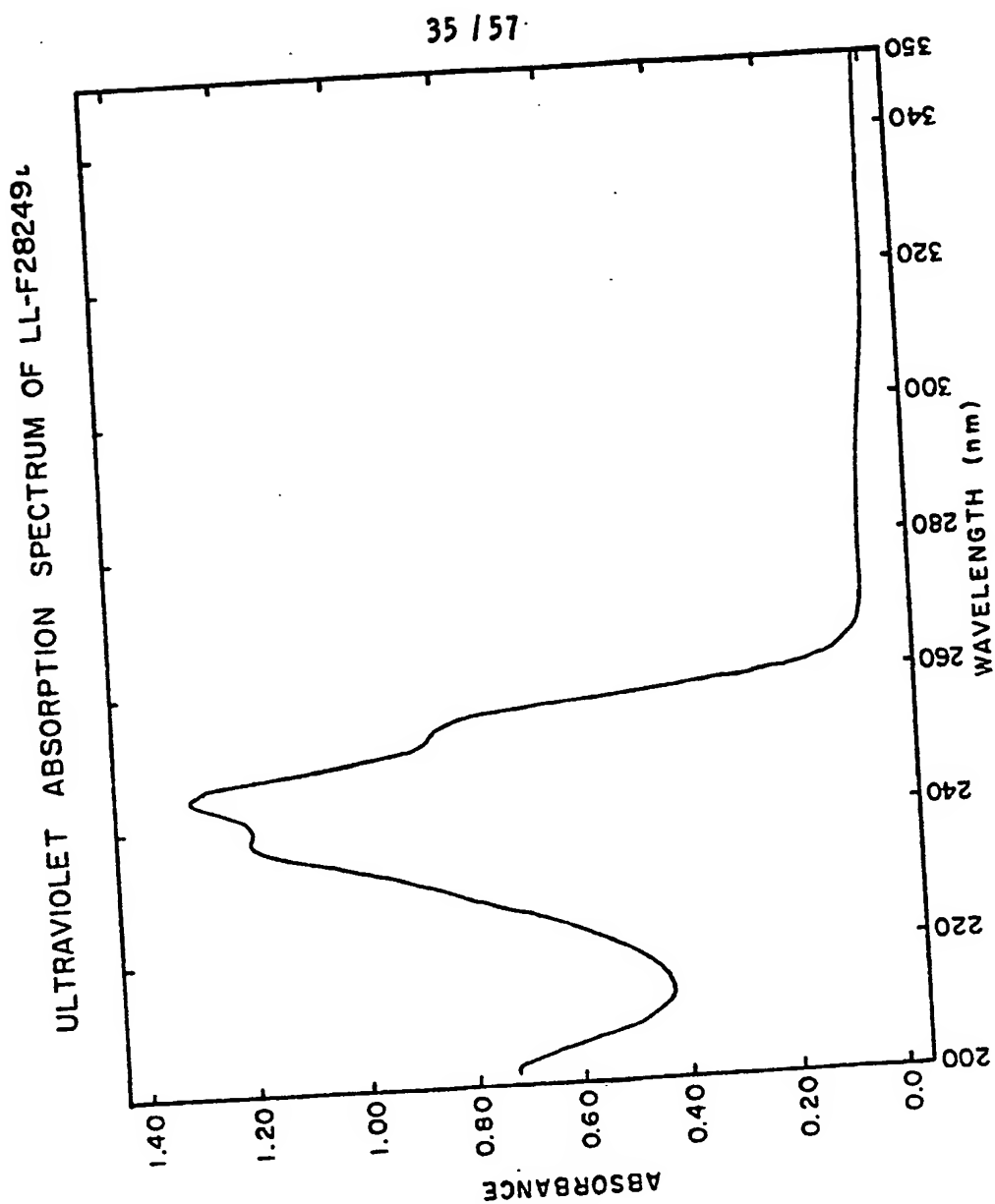


FIGURE 34

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PROTON NUCLEAR MAGNETIC RESONANCE SPECTRUM
OF LL-F28249c IN CDCl₃ SOLUTION

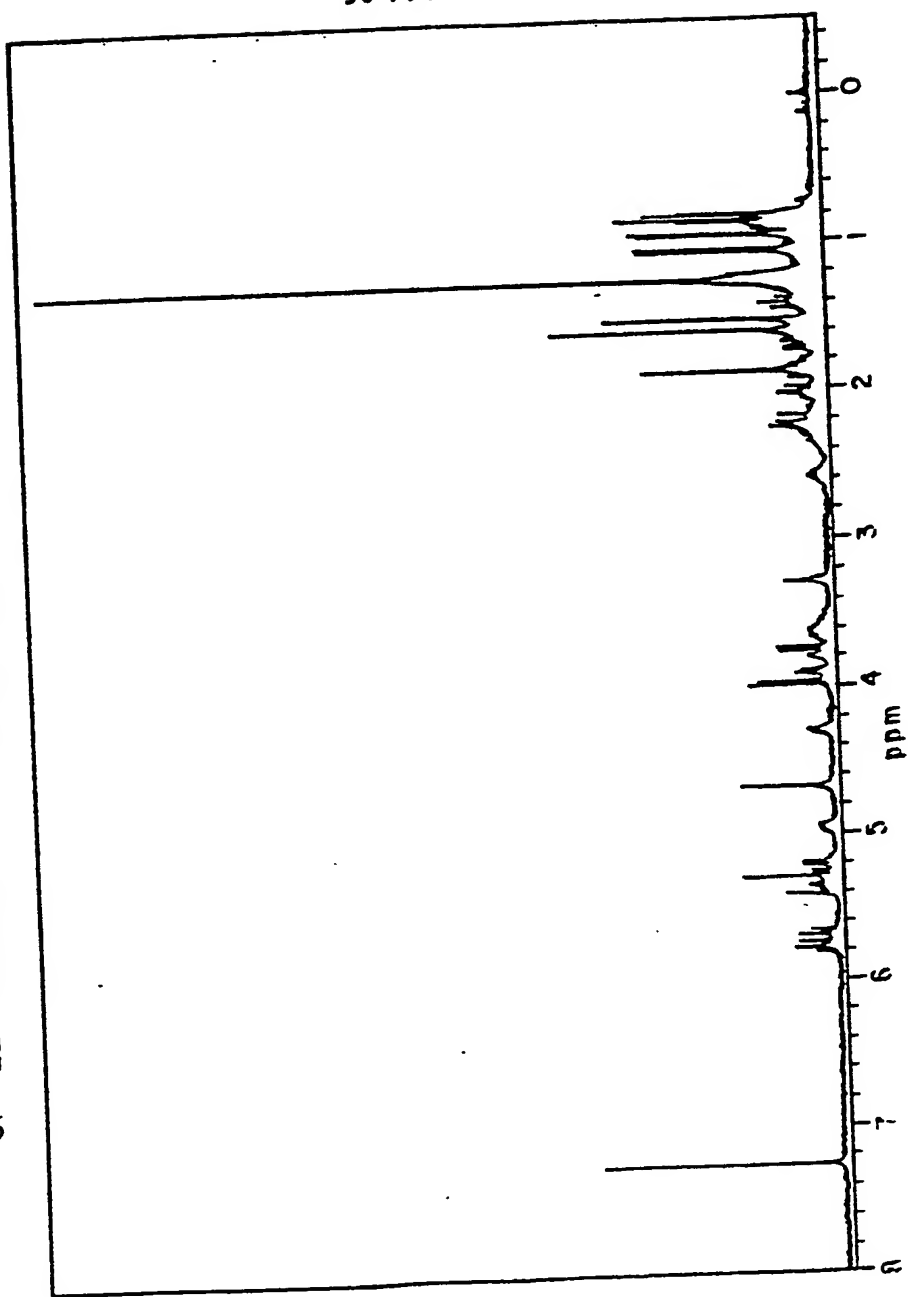


FIGURE 3b

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ELECTRON IMPACT MASS SPECTRUM OF LL-F28249L

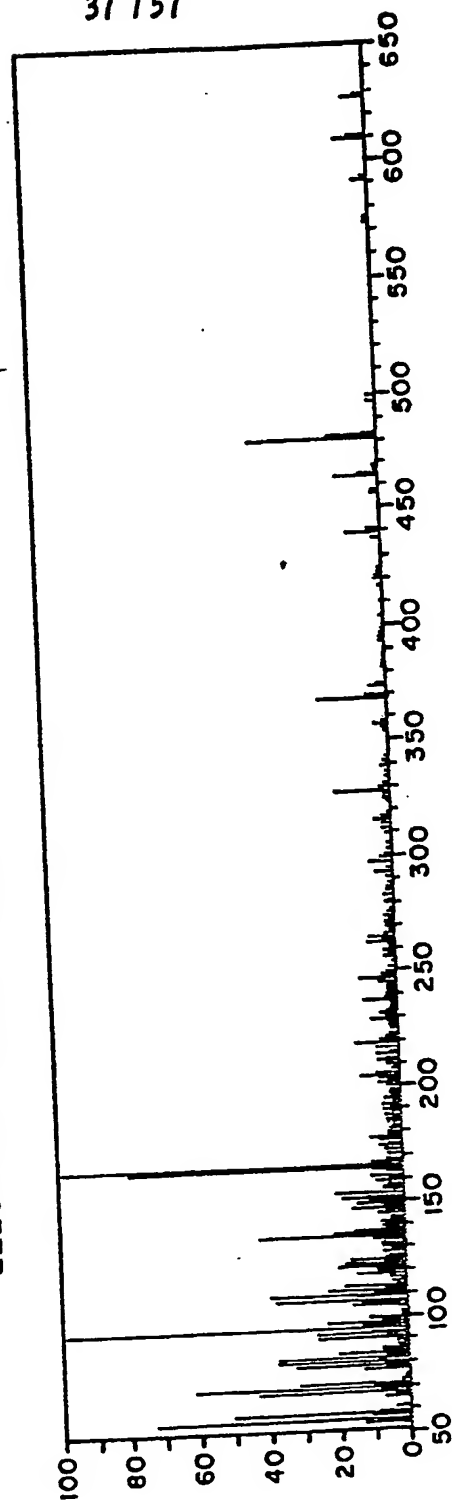


FIGURE 37

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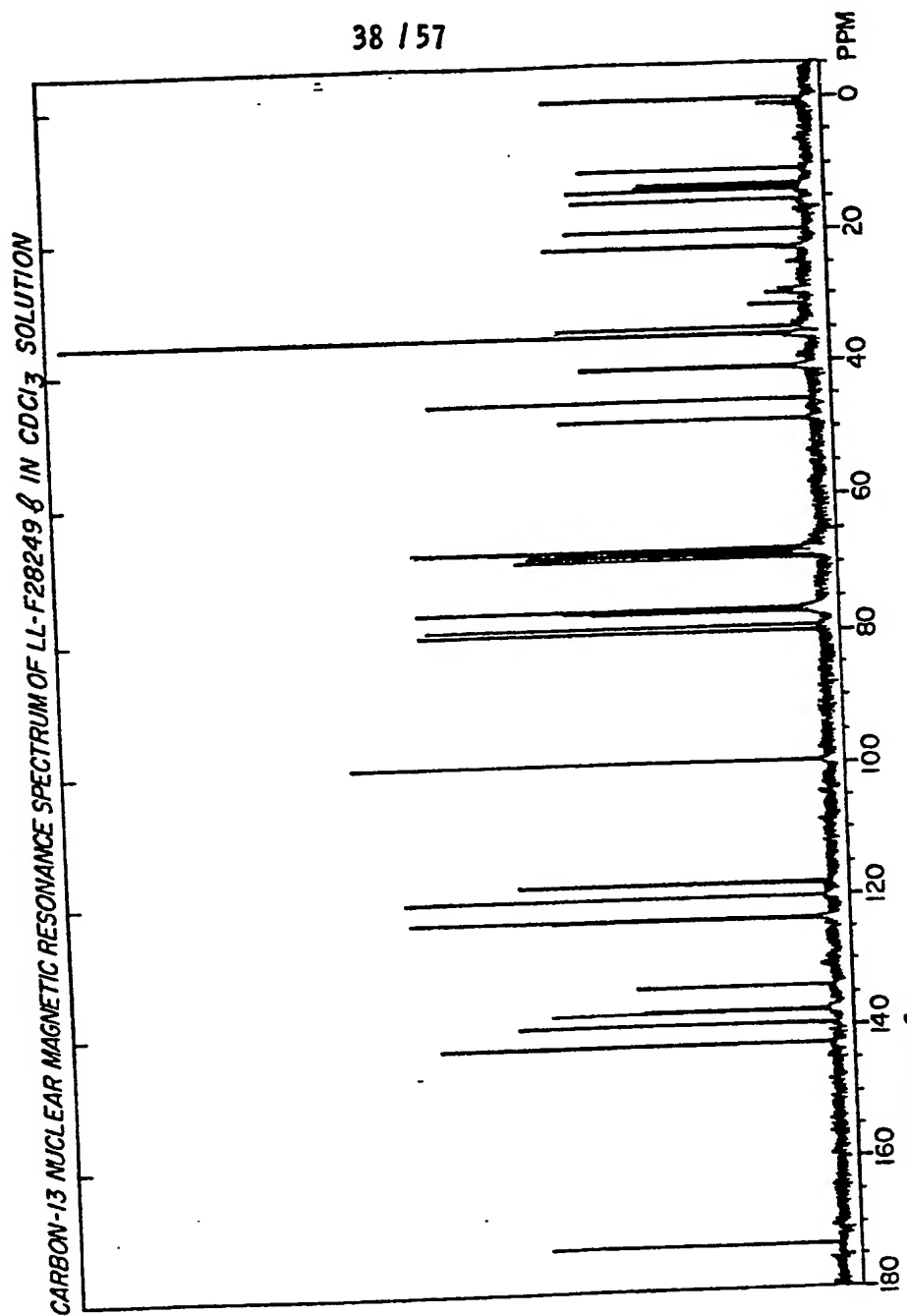


FIGURE 38

0170006

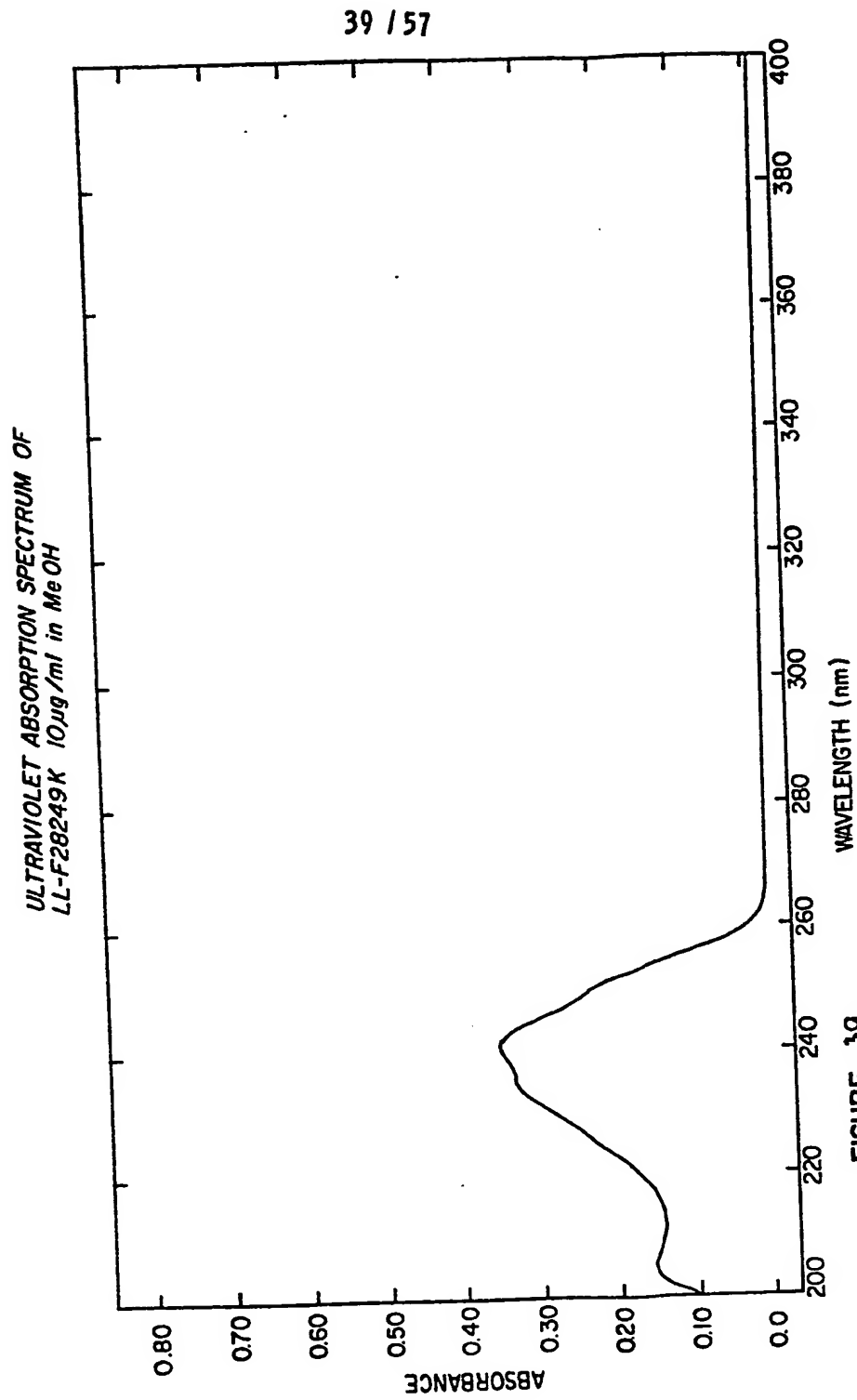


FIGURE 39

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INFRARED ABSORPTION SPECTRUM
OF LL-F28249K (KBr disc)

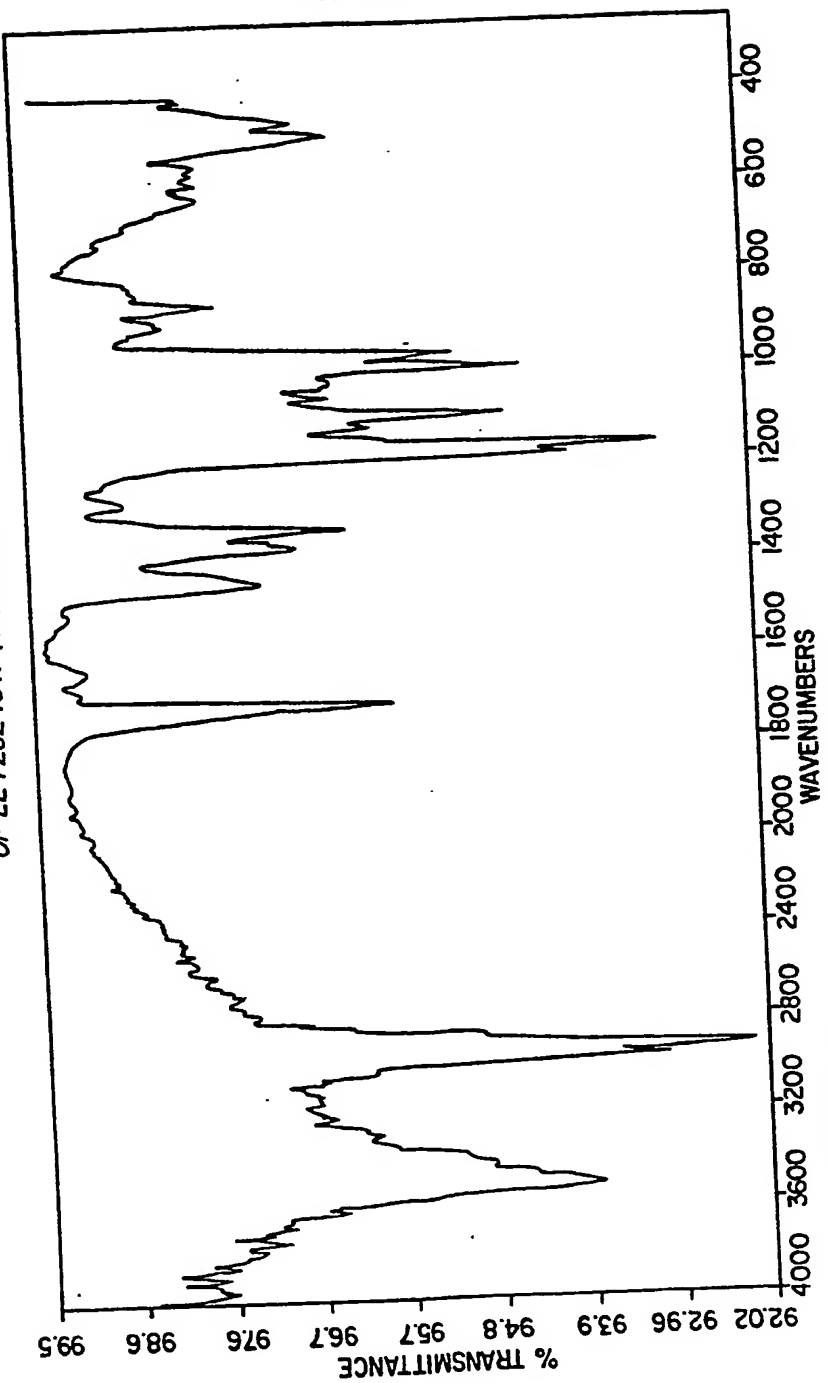


FIGURE 40

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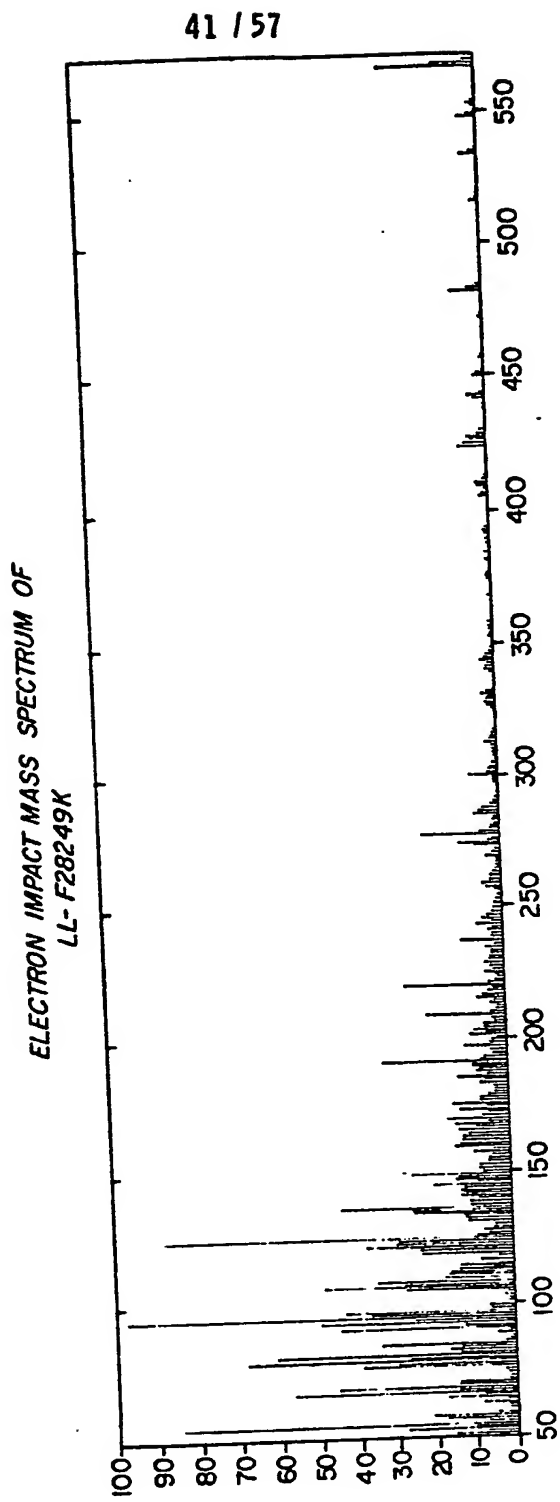
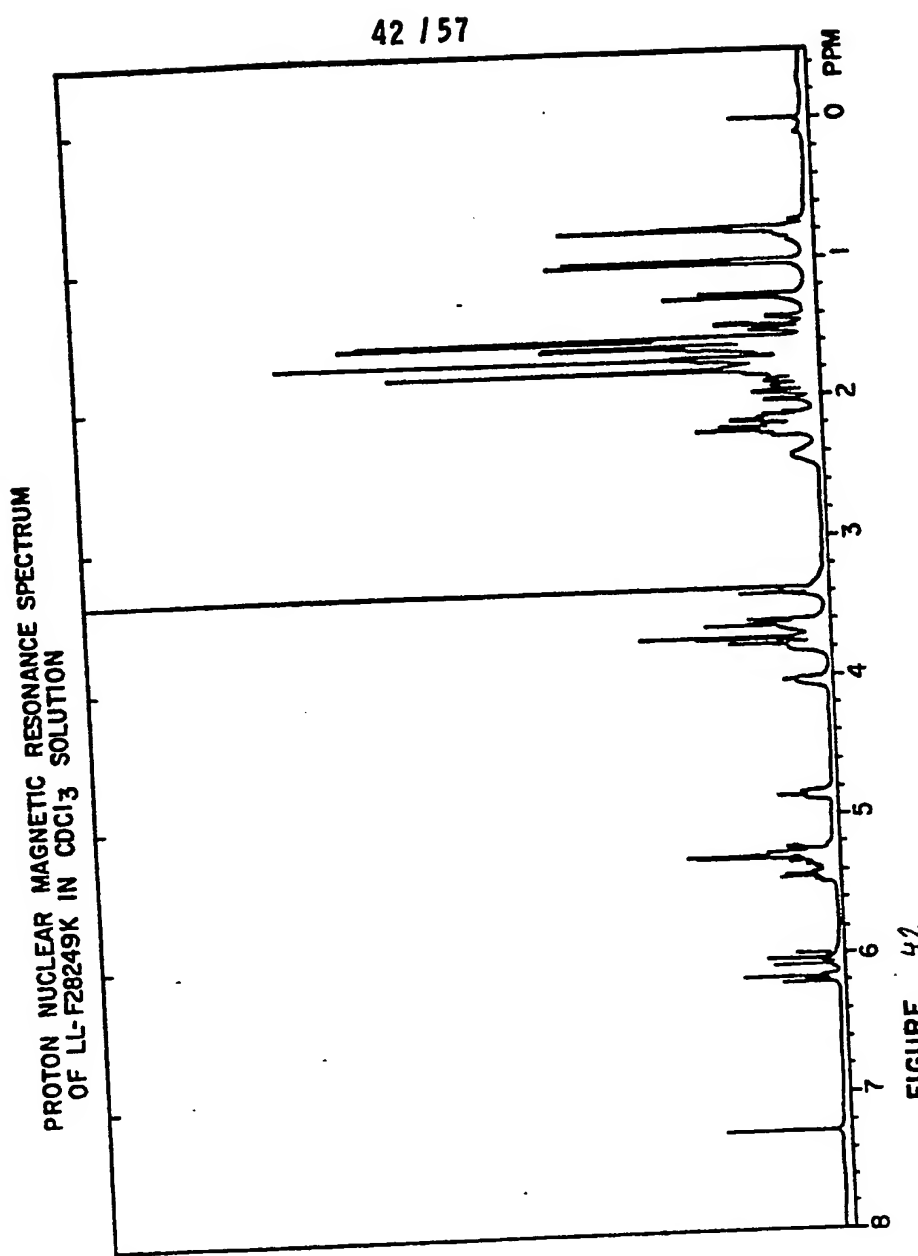
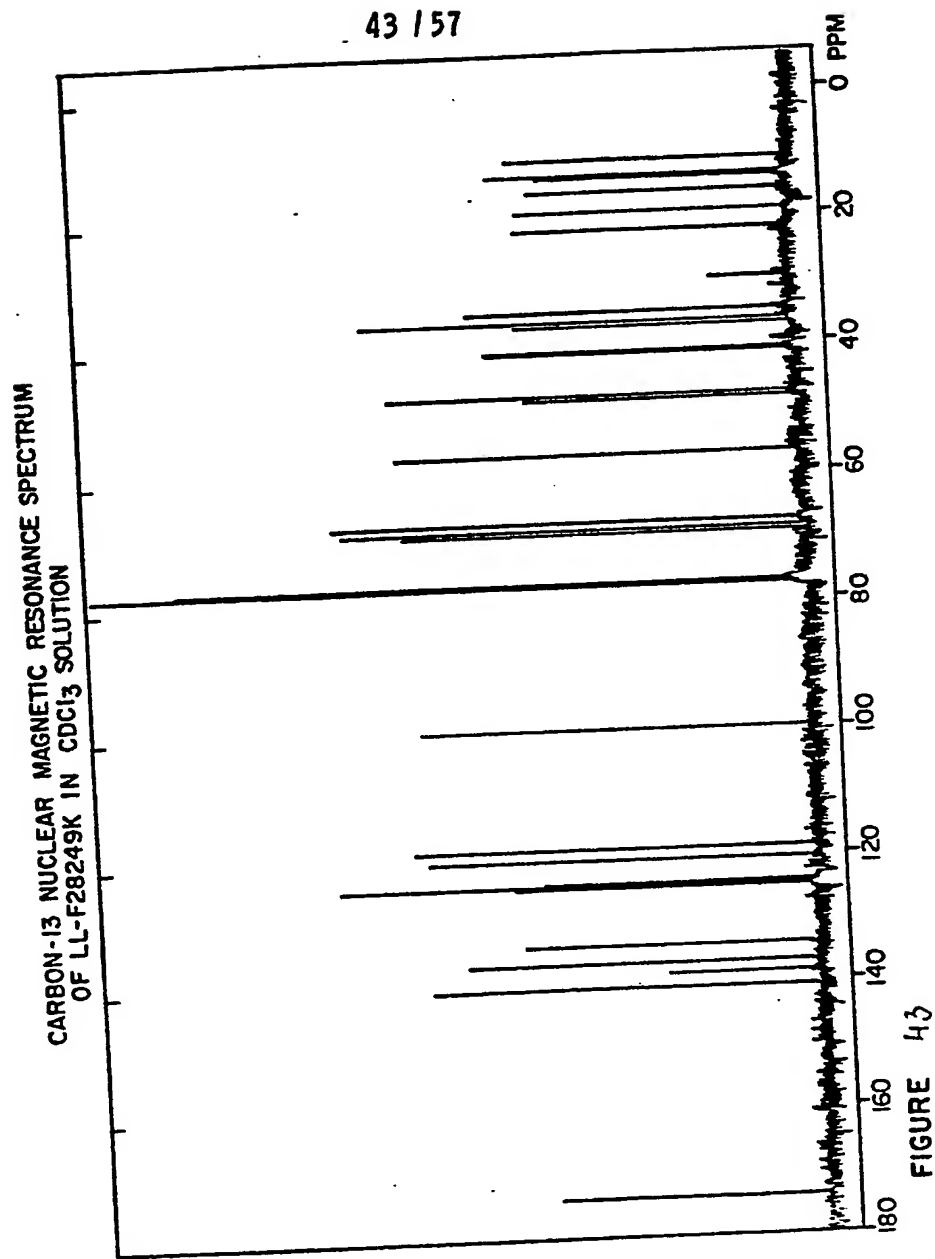


FIGURE 41

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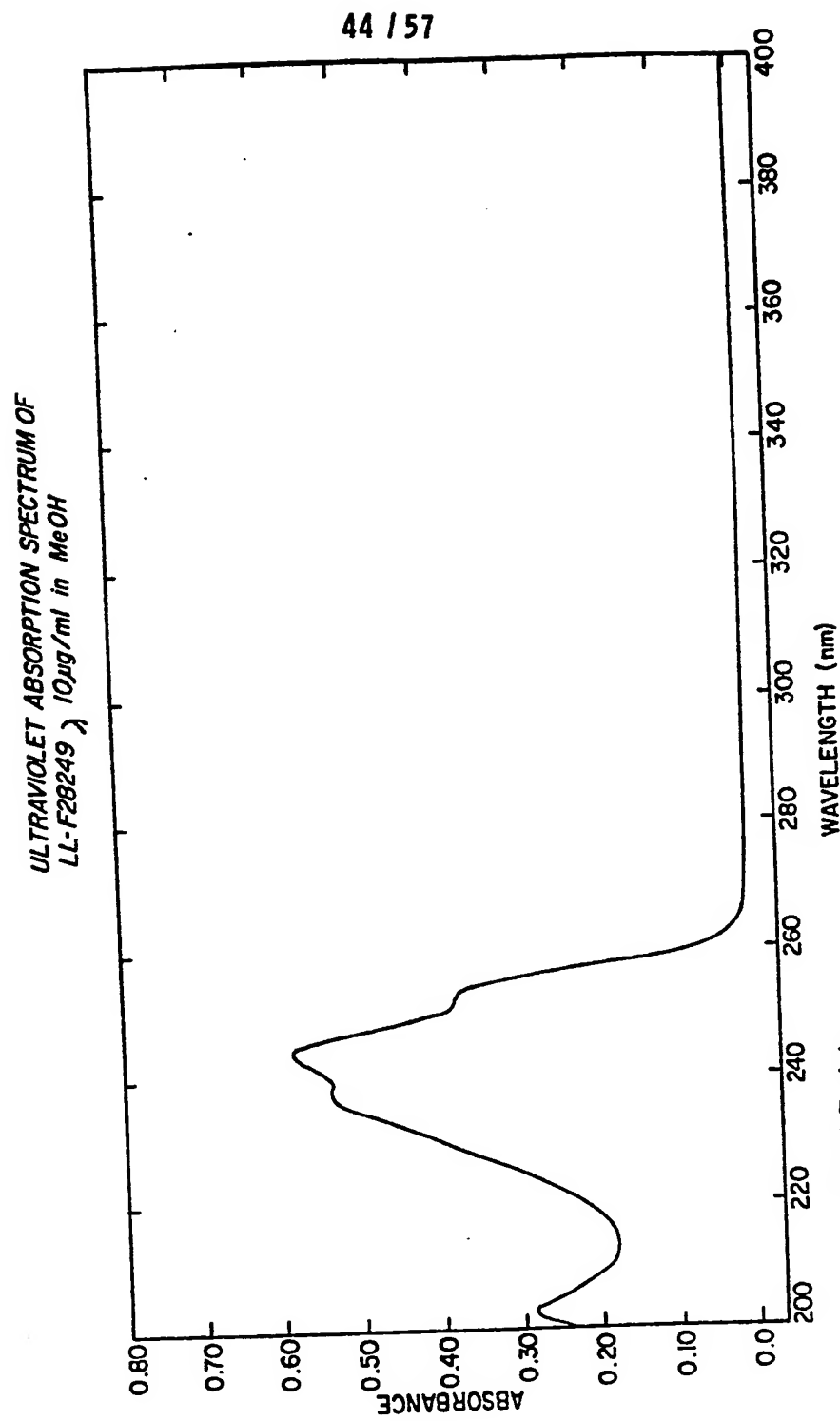


FIGURE 144

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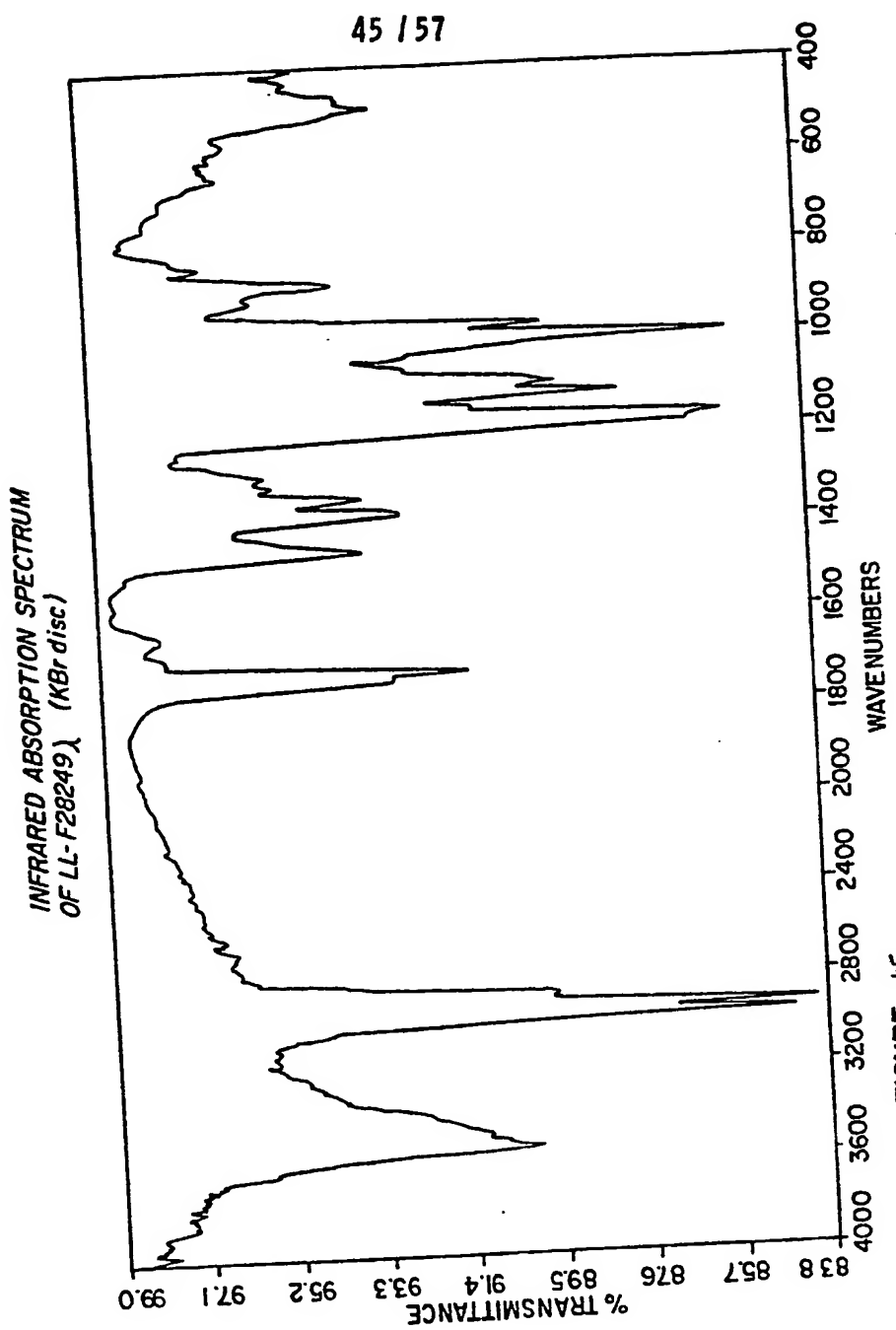


FIGURE 45

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ELECTRON IMPACT MASS SPECTRUM
OF LL-F28249 λ

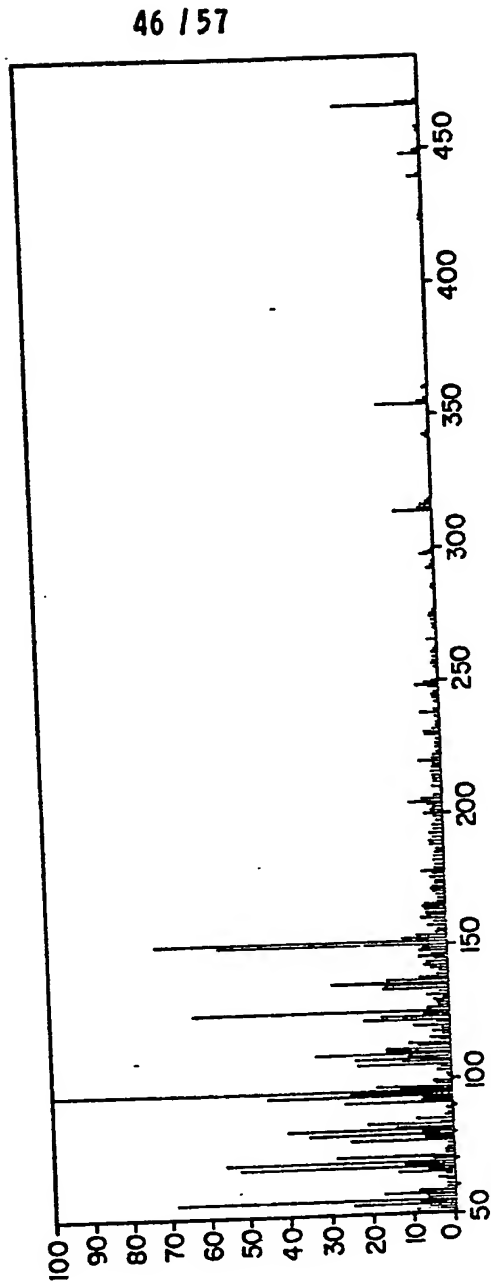


FIGURE . 46

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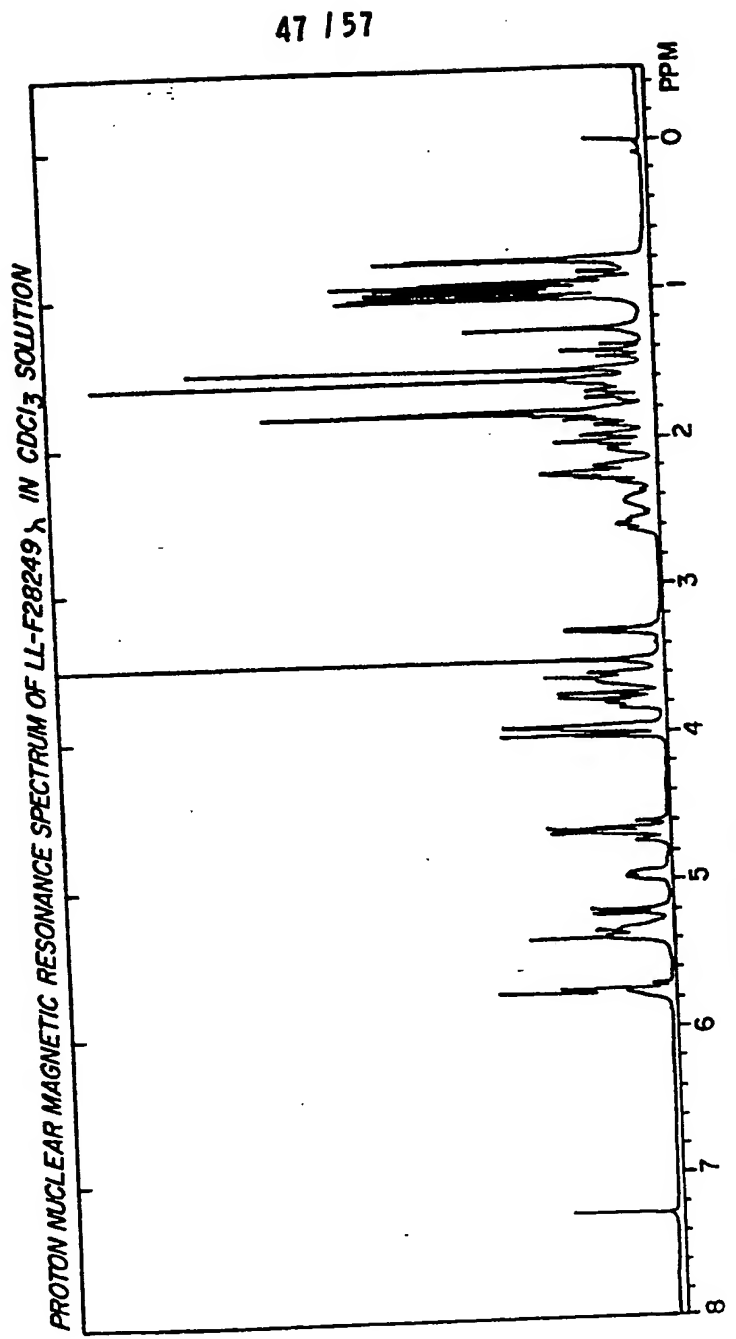
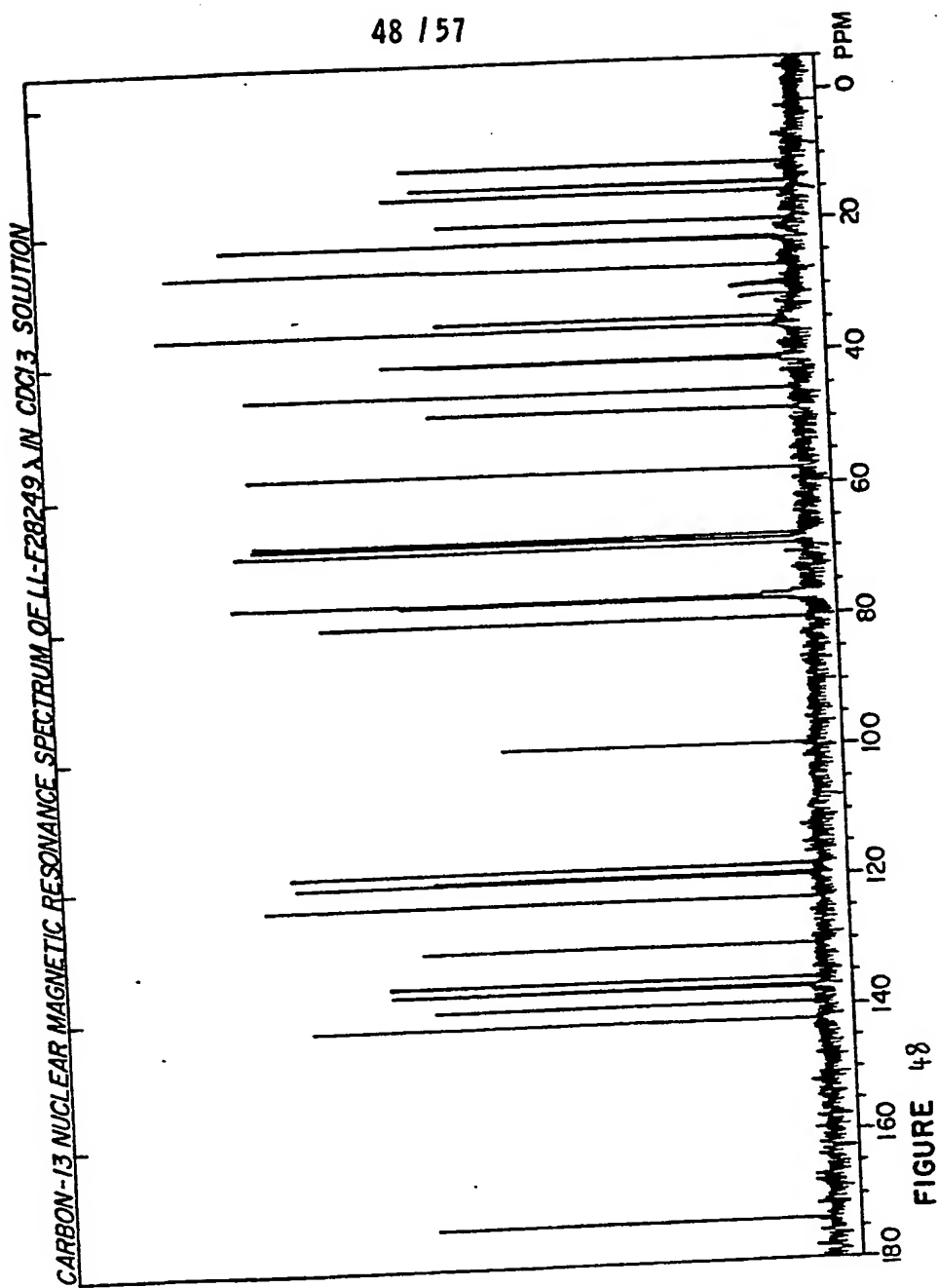


FIGURE 47

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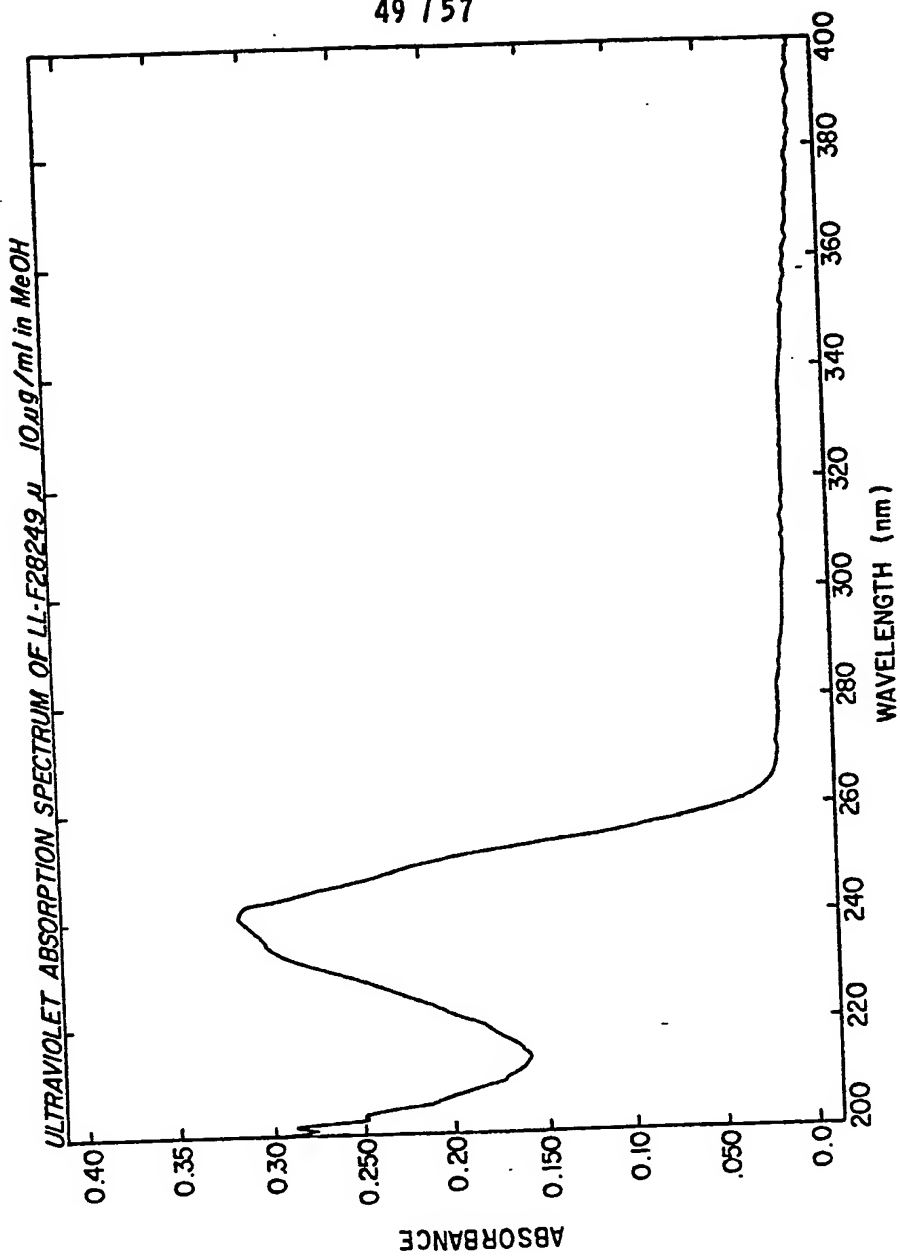


FIGURE 49

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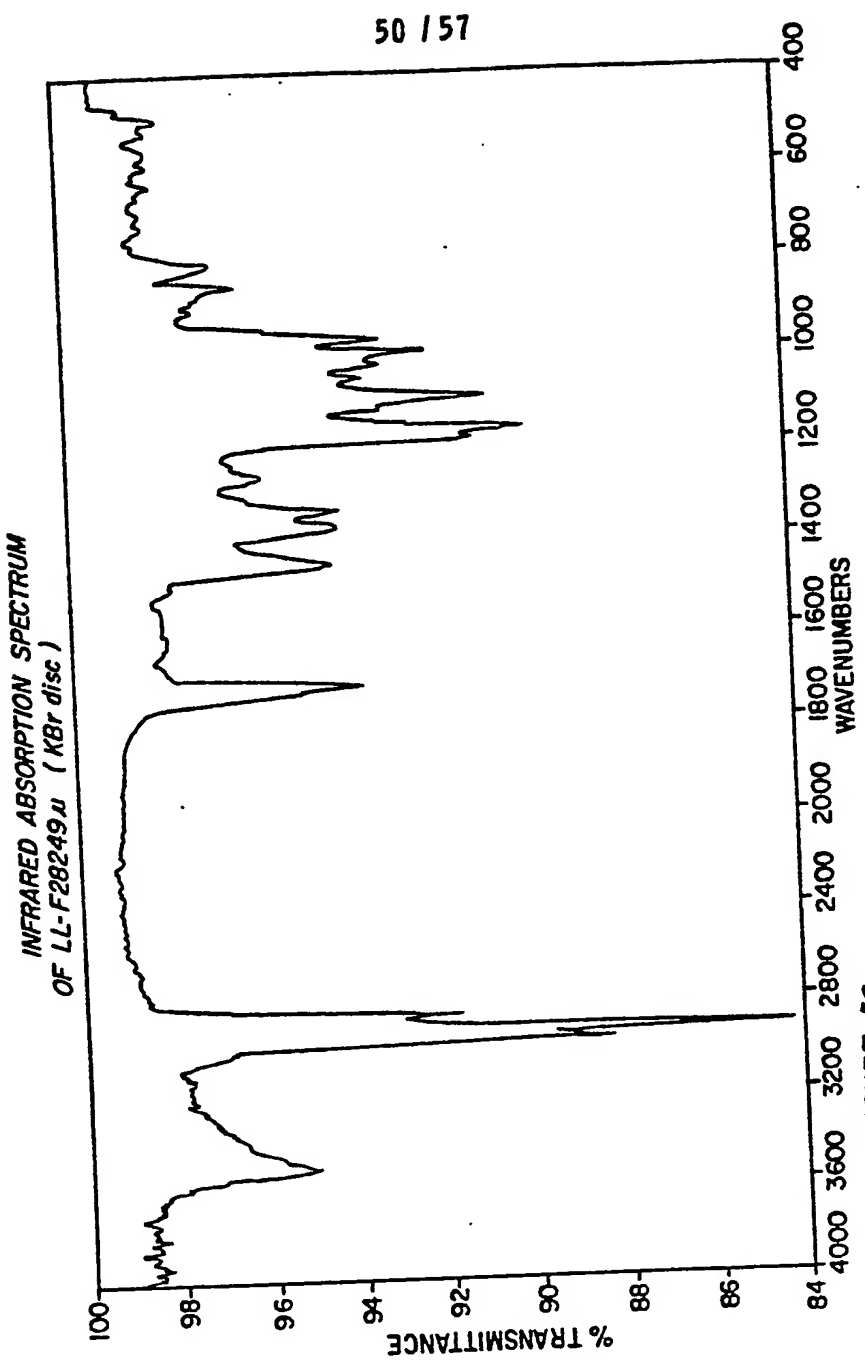


FIGURE 50

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ELECTRON IMPACT MASS SPECTRUM
OF LL-F28249 μ

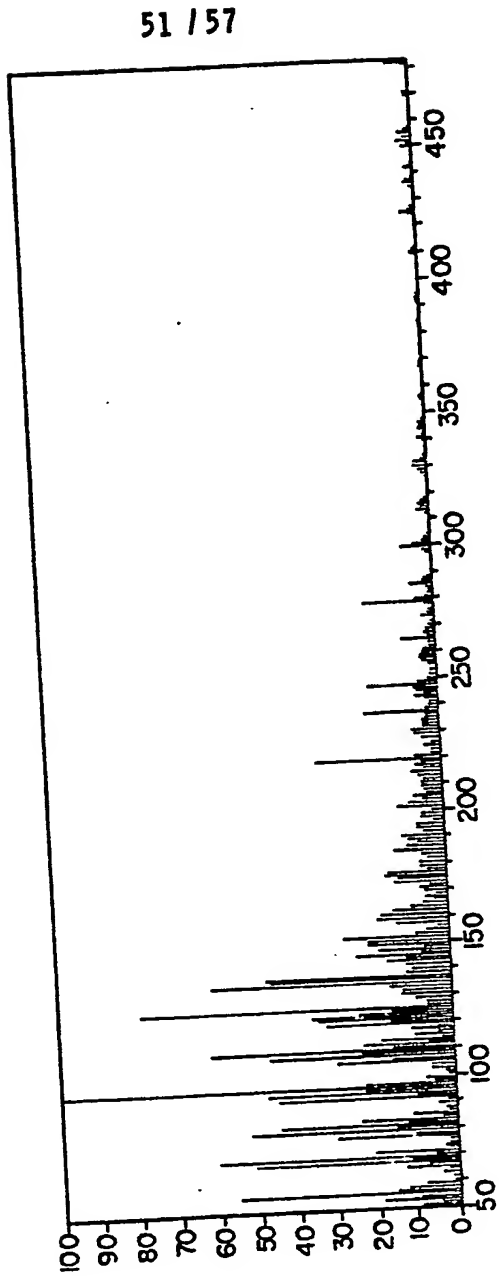
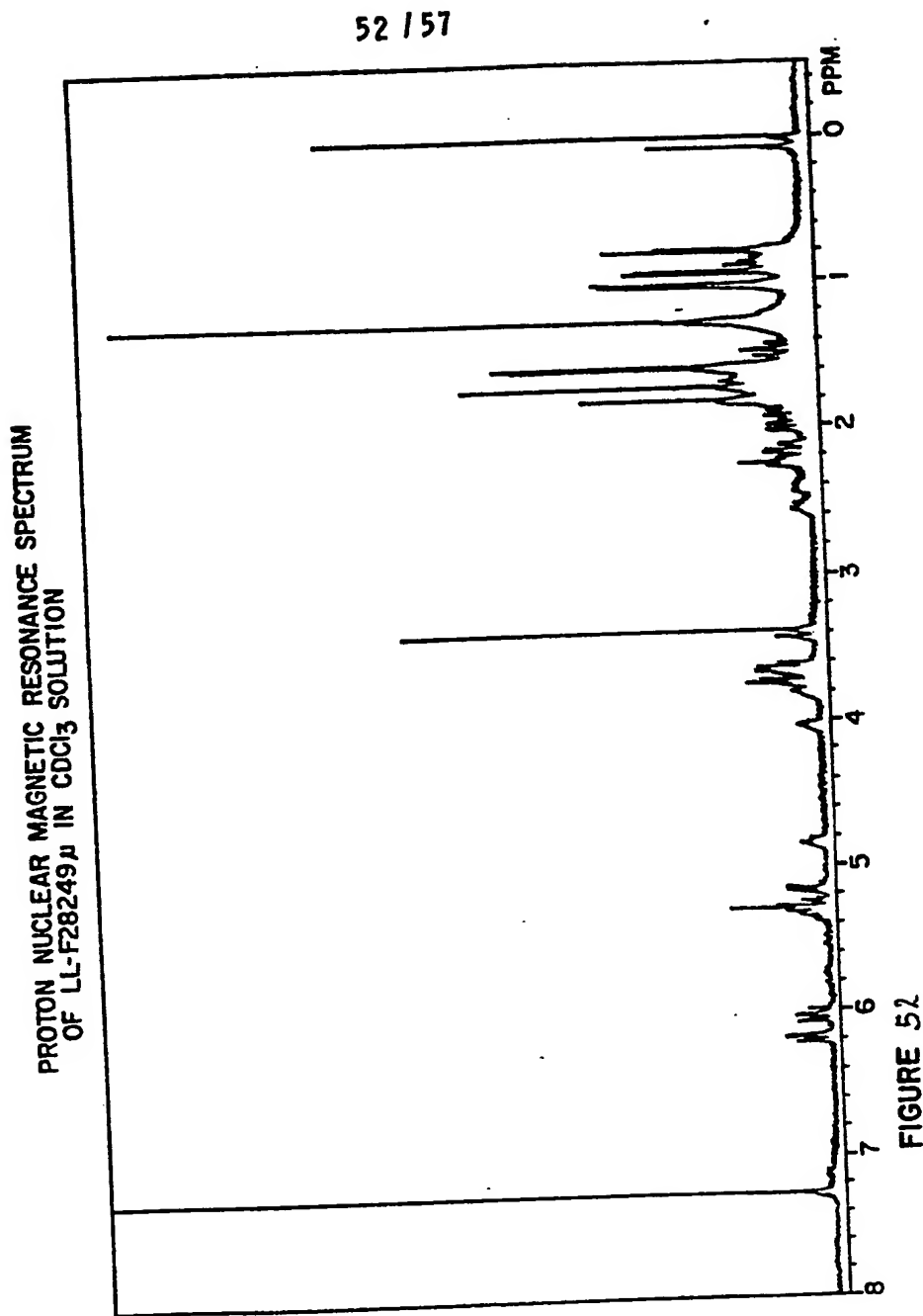
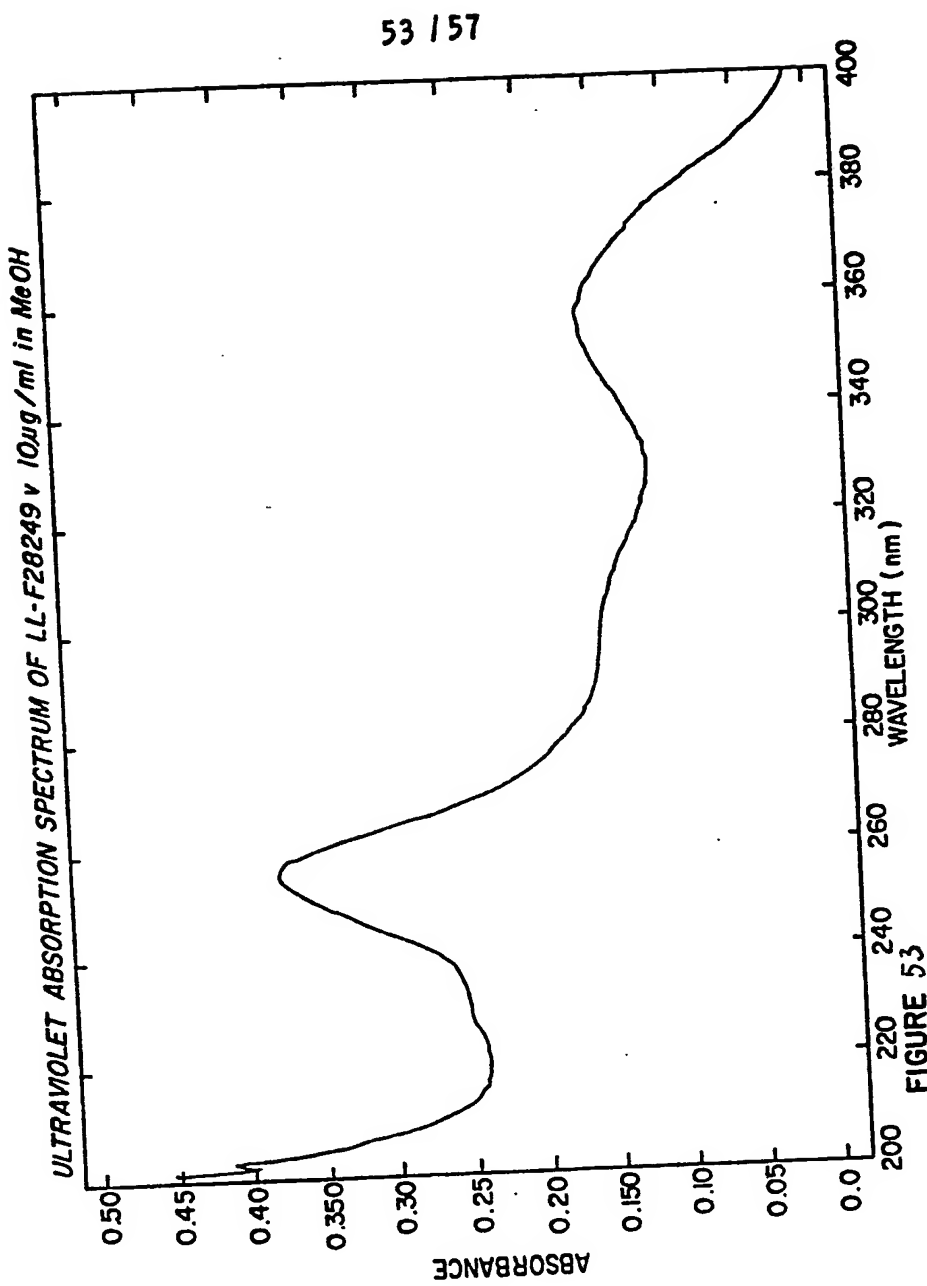


FIGURE 51

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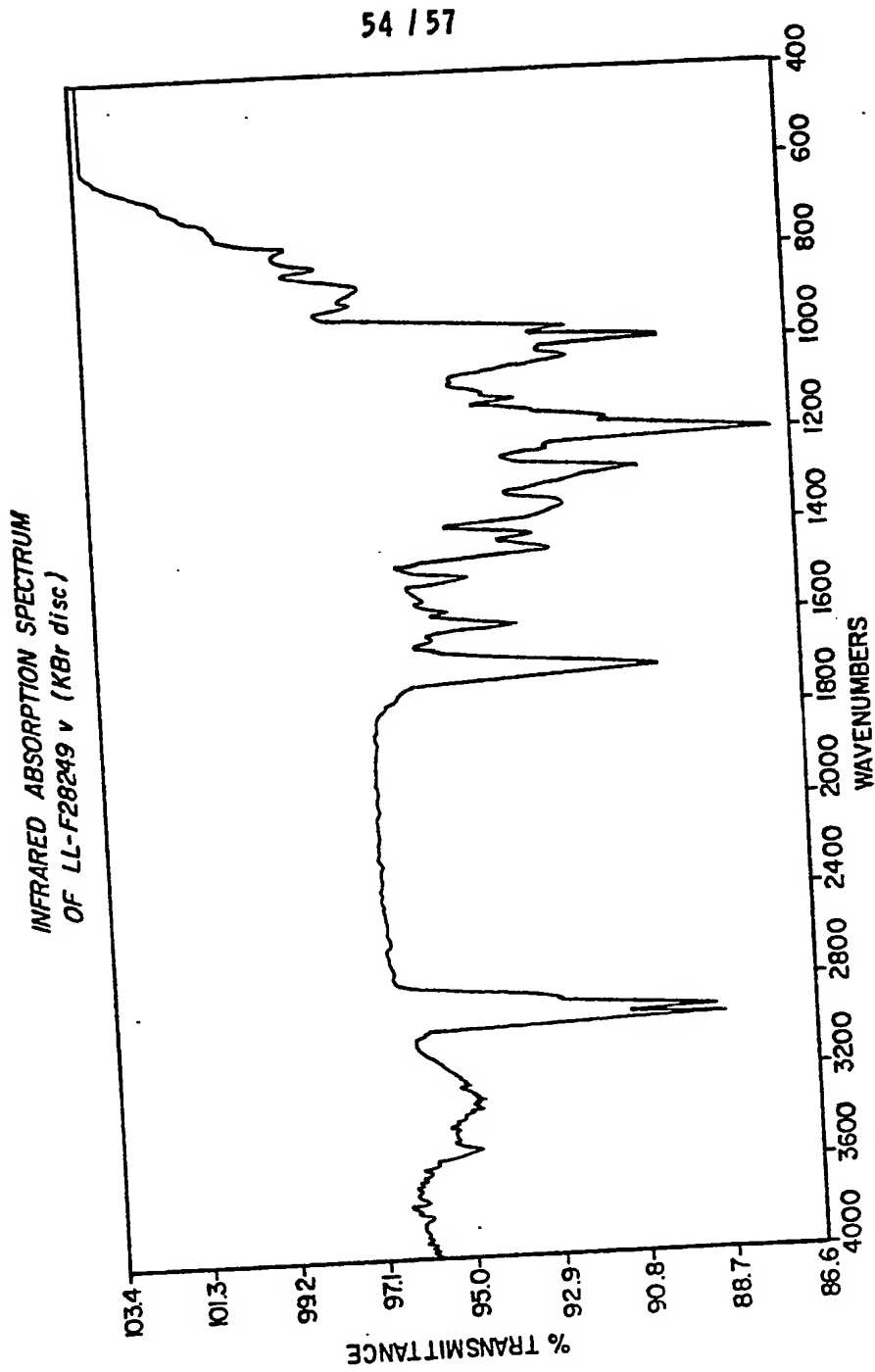


FIGURE 54

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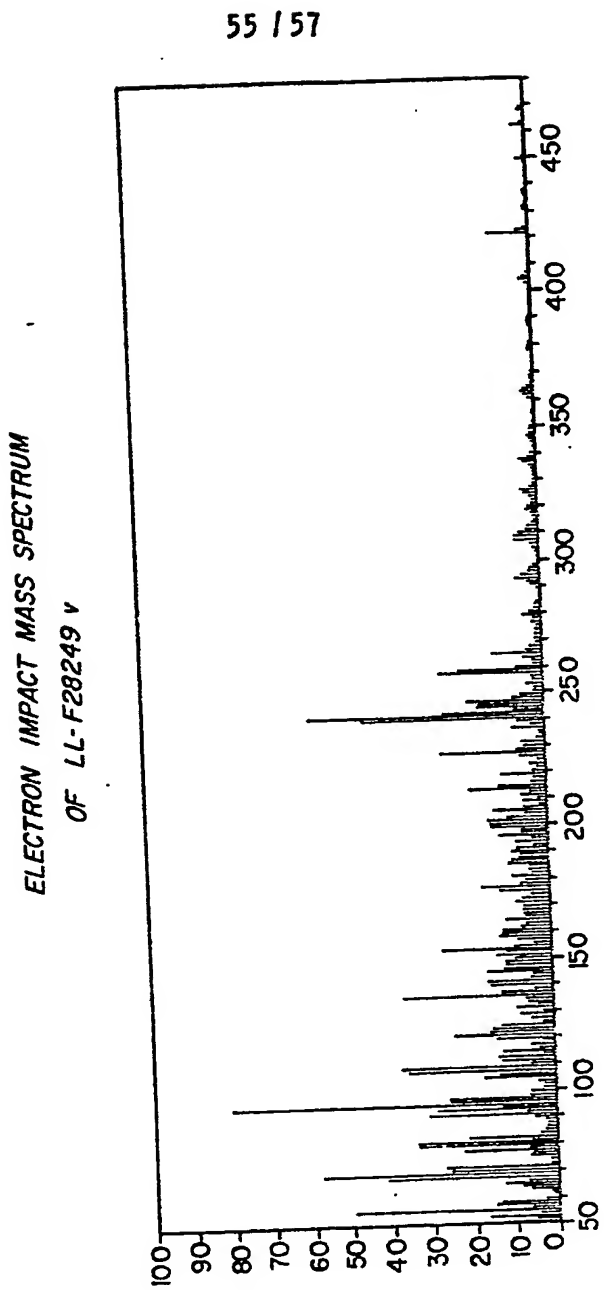
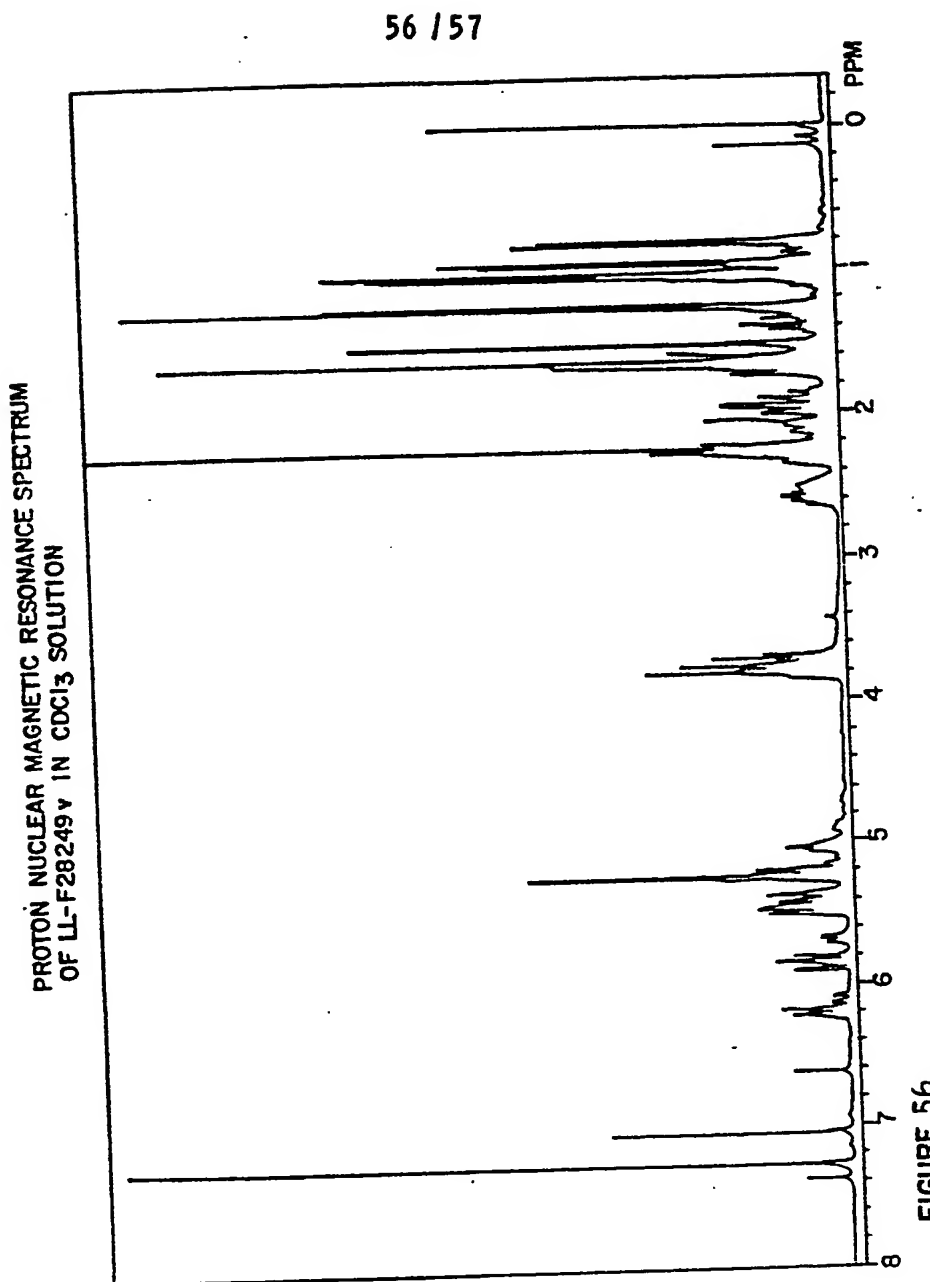


FIGURE 55

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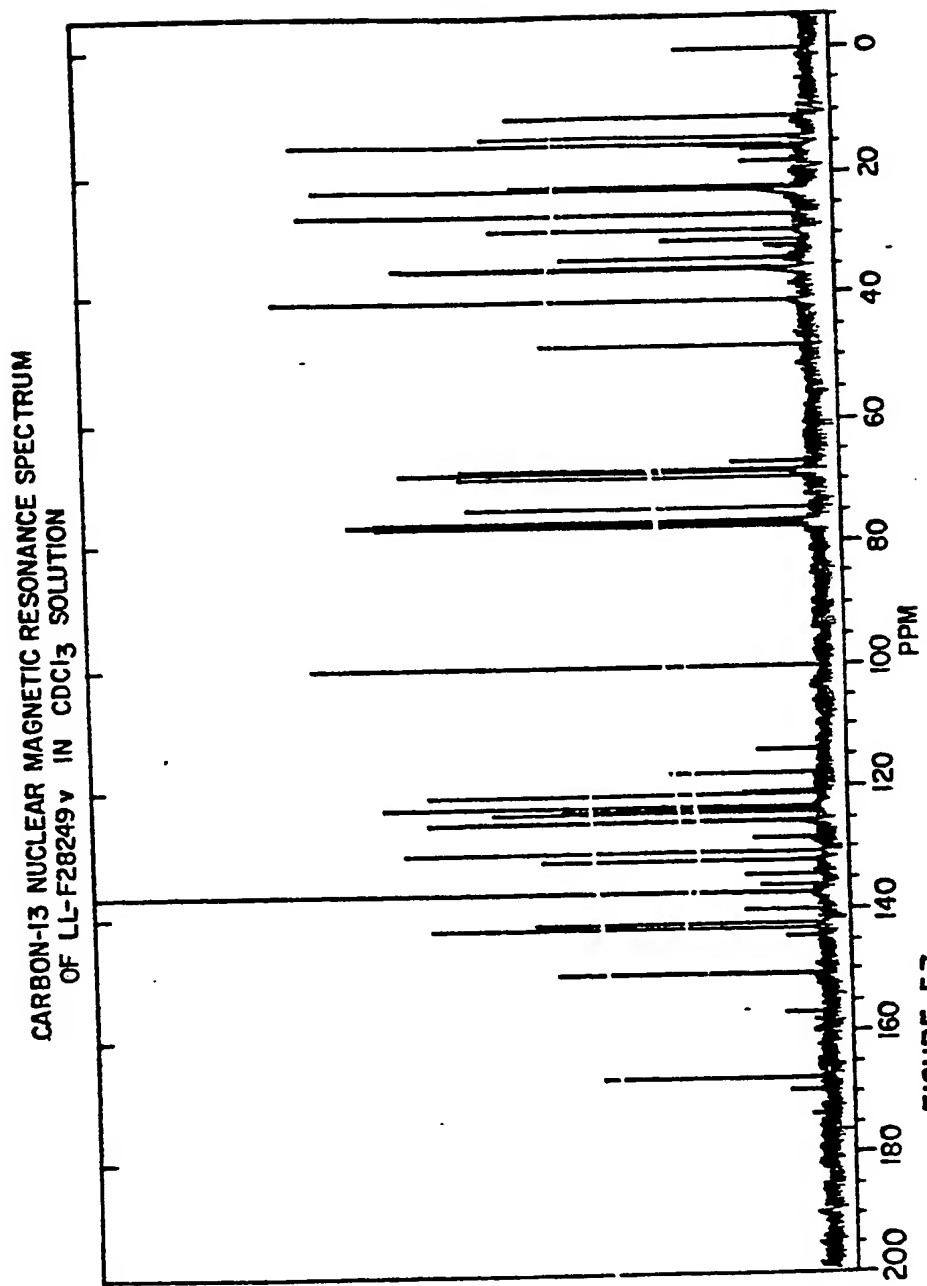


FIGURE 57



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EP 0 170 006 A3

(54) Method and compositions for helminthic, arthropod ectoparasitic and acaridial infections with novel agents.

(55) The present invention relates to novel agents, to their production by fermentation, to methods for their recovery and concentration from crude solutions, to processes for their purification and to pharmaceutically and pharmacologically-acceptable salts thereof. Also, this invention relates to methods and compositions for the control and prevention of helminthic, arthropod ectoparasitic and acaridial infections, in warm-blooded animals, such as meat-producing animals, and poultry, by administering to said animals a therapeutic-

ally or prophylactically-effective amount of new agents designated LL-F28249α, β, γ, δ, ε, ζ, η, θ, ι, κ, λ, μ, ν, and ω or mixtures thereof. The invention also relates to methods for the control of plant nematode infestations and other insecticidal activities. These novel agents are produced via a controlled conditioned microbiological fermentation using *Streptomyces cyanoeigriseus* ssp. *noncyanogenous*, designated LL-F28249 and having deposit accession number NRRL 16773.



EUROPEAN SEARCH REPORT

EP 85 10 6844

DOCUMENTS CONSIDERED TO BE RELEVANT			CLASSIFICATION OF THE APPLICATION (Int. CL4)
Category	Citation of document with indication, where appropriate, of relevant passages	Relevant to claim	(Int. CL4)
Y	EP-A-0 058 518 (MERCK) * Whole document * ---	1-14,20	C 12 P 1/06 C 07 D 493/22 C 07 G 11/00 A 61 K 35/66 A 61 K 31/35 C 12 N 1/20 A 01 N 63/02 A 23 K 1/17 // (C 12 P 1/06 C 12 R 1:465 C 12 N 1:20 C 12 R 1:465) (C 07 D 493/22 C 07 D 313:00 C 07 D 311:00 C 07 D 311:00 C 07 D 307:00)
Y	EP-A-0 102 721 (SANKYO) * Whole document * ---	1-14,20	
Y	CHEMICAL ABSTRACTS, vol. 85, 1976, page 196, abstract no. 118436e, Columbus, Ohio, US; G.T. CARTER: "I. Structures of oligomycin A and C. II. Structures of three isomeric octadecadienoic acids possessing divalent cation ionophoretic activity. III. Insecticidal components of dill and anise plants", & DISS. ABSTR. INT. B 1976, 37(2), 766-7 * Abstract * ---	1-14,20	
Y	JOURNAL OF ANTIBIOTICS, vol. 36, no. 8, August 1983, pages 980-984, Tokyo, JP; H. MISHIMA et al.: "Milbemycins, a new family of macrolide antibiotics structure determination of milbemycins D,E,F,G,H,J and K" * Whole article * ---	1-14,20	TECHNICAL FIELDS SEARCHED (Int. CL4)
Y	EP-A-0 073 660 (MERCK) * Whole document * ---	1-14,20	C 12 P
L	CHEMICAL ABSTRACTS, vol. 106, 1987, page 511, abstract no. 32663s, Columbus, Ohio, US; G.T. CARTER: "Structure determination of oligomycins A and C", & J. ORG. CHEM. 1986, 51(22), 4264-71 * Abstract * -----	1-14,20	
The present search report has been drawn up for all claims			
Place of search	Date of completion of the search	Examiner	
THE HAGUE	17-03-1988	RAJIC M.	
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